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## SYSTEMATIC QUALITATIVE ORGANIC ANALYSIS

BY

H. MIDDLETON

MSc. (Manchetta), A.I.C.

LECTURES IN ORGANIC CHEMISTRY.

FRADPORD TECHNICAL COLLEGE



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#### PREFACE

This book is intended mainly for the training of students in the general methods employed in the identification of organic comrounds.

During the preparation of the book the author personally examined over 600 purely organic substances and a large number of their metallic derivatives. Conditions, necessary for a definite result, were found for every test, and then systematic schemes of analysis were built up. The solid derivatives, to be prepared for identification purposes, have been carefully arlected. Where alternatives were possible, those with very low or very high melting points have been avoided. The author has prepared all the derivatives mentioned and determined their melting points. This work has been carried out in order that detailed methods of preparation. either general or special, could be given; also in a considerable number of cases for the purpose of deciding between the widely different values for the melting point of a particular compound given in works of reference. For rurnoses of economy, derivatives for whose preparation only relatively inexpensive reagents are required, have been chosen.

By following out the schemes and the instructions for the perparation of derivatives students obtain decisive results, and

thus gain confidence in their work.

Particular attention has been paul to the requirements of pharma-

Introduct attention has been paul to the requirements of pharmacentical students, the schemes permitting of the rapel identification of the majority of the single organic compounds included in the Petitah Pharmacoporia.

The time factor has always been kept in minf, the minnim times for the various transitions having been found. The times required for the hydricipus, under the specified conditions, of always 100 setters are pires. Thus if an esser is required to be hydricipus in an examination, or during a levil practical period, one which is extilable may be chosen from the fact.

For students taking lecture course, but not requiring analytical work, the head provides rapid exercises in the provision explicited in presental explain work. Using the small quantities of substances indicated, even part time day or evening students can pain considerable tractical experience in a short time, and w

Little expense to the institution they are attending. A specia of such exercises, and involving analysis, is included,

Certain classes of compounds have been emitted, as it sidered unlikely that students will encounter them; al blentification of dyes and their more complex intermedi-

considered beyond the sease of the work. The author has aimed at producing an executially practical hence chemical equations and explanations have been or Students of Organic Chemistry will usually be attending ! courses, where equations will be given and reactions expl In the case of the sumpler compounds given for klentification suggested that, after the completion of their practical work, str should give an account of the reactions involved, gaining required information from lecture notes, text-books, or sources. Knowledge gained in this way will be far more int ing, and will be retained for a longer period, than if acq without effort.

As but few names are mentioned, a general acknowledgme here made to all chemists whose work has been utilised.

The author is greatly indebted to his colleague, Mr. M. G. wood, M.A. (Oxon.), for his careful and helpful criticism of the

H. MIDDLETO

THE SENIOR CHEMICAL LABORATORY. BRADFORD TECHNICAL COLLEGE. December 1938.

#### PREFACE TO THE SECOND EDITION

Some of the schemes for the determination of the class o compound have been rewritten so that by applying all the tests the scheme considerable information concerning the presence groups in an organic compound, not listed in the book, may gained.

About thirty more compounds, mainly aldehydes and keton have been added to the lists of substances.

H. M

September 1943.

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#### STANDARD ABBREVIATIONS

	regent	١.								aqueor
b.p.										boiling poin
conc.										concentrate
0.0.									cubic	centimetre
d. (ale	TAVE (	ollav	red by	6 D	umber	ı .				densit
							ind	icates an	annr	oxi-
			e norn							. dilut
quiv.	·									equivalen
ğ			-							. gram(a
n.p.						·			172	elting point
	conne	etior	ı with	etan	dard a	olotic	ons)			. normal
			with						•	. norma
pt.										precipitate
ec.	·	•	•		•					secondary
oln.	Ť	Ť	Ĩ.	Ť	Ţ.	Ť	Ċ			solution
ert.	·	•			·	:	Ċ			tertiary
.t	Ċ	•	•		•	:	Ċ			tost-tube
ol	Ċ	•		Ť		Ť				. vojume
rt.	•	•	Ċ	÷		:	Ċ			weight
ecomp	lin e	onn.	ection	with	m.n.l			wit	h dec	mposition
[o .							Ċ		Moth	yl, CH.
t .			-						Ethy	1, C,H.
ъ.	:			·	Ċ				Phon	yl C,H,
								Ac	etyl,	CH, CO-
ı.								Benz	oyi, C	HCO
	:	:						Radical	(alky	or aryl)
•	•	•	•							
			SPEC	IAI,	ABBR	EVL	ATIO		rinal a	ubetance
в.	•	•	•	•						oo grain
a.			•	•			-			-

# SYSTEMATIC QUALITATIVE ORGANIC ANALYSIS

# INTRODUCTION

canic substances are carbon compounds,

presence of carbon in a substance is indicated by one or or two somewing resume.

The liberation of carbon, which may be left as a black residue

when the substance is hested on platinum or porcelain, or in the case of inflammable substances deposited as soot when in the case of minimum and substances dejended as much when a cool surface is held in contact with the flame of the burning mbetance.

In either case the black substance is proved to be carbon by the fact that on heating to redness it burns away com-

he production of a characteristic blue flame, due to the urning of carbon monoxide, when the substance is warmed triang or careful mith concentrated sulphuric acid and the mouth of the tube turned to the flame.

(3) The milkiness produced in himewater by passing through it

(a) a dilute mineral acid is added to the substance. This applies to most carbonates, those containing a

metal being usually regarded as inorganic substances. (b) a mixture of the substance and finely divided copper oxide is heated to redness, or the rapour of the substance is passed over red-bot copper oxide in a suitable

For the identification of a single carbon compound it is greerally adrivable first to test for the elements present.

It is assumed that the substance under examination will contain as a more of the following elements: hydragen, outpen, nitrogen, sulphin, chlorine, bromme, kodine, phosphoren,

The test for hydrogen, which depends on the formation of water (detected by the blue colour firm to achydrous copper Before recursively work the introduction about he carefully read.

sulphate) when the substance is heated with copper coids is not worth while, owing to the time involved in the complete drying of the substance and the copper coids. The few common right substances which, if anhydrous, contain no hydrogen (e.g. carbon tetrachloride, herarchiorocthane, metallic oxalates, etc.) present no difficulty in defection.

The presence of exygen is usually inferred by the recognition, during the kientification, of a characteristic group e.g. —OH.

The next procedure is usually the determination of the class of the compound, i.e. to ascertain whether it is an alcohol, earbohydrate, amine, etc.

Finally the compound is identified as a particular member of a class by some or all of the following methods:

(a) comparison of the physical properties of the substance with those of known members of the class.

(6) application of special chemical tests.

(c) preparation from the compound, if possible, of a pure solid substance known as a "derivative," and determination of its melting point. Reference is then made to a list of melting points of the particular derivatives in order to ascertain the melting point obtained is identical with, or near to, that of one of the derivatives in the list.

(d) quantitative work, e.g. determination of the equivalent weight of an acid, or estimation of the percentage weight of a par-

ticular group present.

#### EXPERIMENTAL TECHNIQUE

The substance to be identified will be referred to throughout the book by the letters O.S. (= original substance).

(1) Measurement of quantities, etc.

In all the tests given the quantities which produce suitable results have been found by trial, hence to ensure equally definite results quantities approximately equal to those mentioned should be used (the word "approximately" should not be interpreted too filterally). For either a solid or a liquid a certain depth (referred to for convenience as a "layer") in a test-tube of the width creatly employed (i) in diameter) is mentioned, the depth referring to the vertical distance between the surface of the substance and the lowest part of the curved bottom of the tube

Except when the word "measured" precedes the figure, the measurements are intended to be made roughly with the eye, but it might be advisable at the beginning to glance at a rule.

It should be noted how much of a solid substance it is necessary

to take up on the end of a spatula to approximate to a 1-in. layer in the test-tube mentioned, since it is not to be expected that a dry tube will be used every time and if the tube is wet most of the any sume was no used creek sides and all idea of measurement by

Unless a solid substance is in the form of a reasonably dense powder it should be ground up in a mortar.

In the case of solids the letters R.G are used to indicate an amount roughly equal in bulk to a large rice grain.

By a "trace" is meant an amount roughly equal in bulk to an ordinary pin-head

Volumes of liquids are also referred to in c.c. A.5-in. X f-in. test-tube has a capacity of approximately 20 c.c., hence if this length of tube is employed 5 cc. will refer to a quarter of a test-tube full, and so on. For 1 c.c., 2 c.c., and 3 c.c. it will

to a secretary run, and so on. And a con, and a con, and be sufficient to employ respectively a depth of \$\frac{1}{2}\$ in, \$\frac{1}{2}\$ in, and \$\frac{1}{2}\$ in. o summents we employ respectively a negative 2 mm, 1 mm, and 2 mm. In the tests definite volumes are stated, but they are only intended to be approximate unless in italic type; the same applies to the (2) Use of litmus paper.

Experience has shown that often correct results are not obtained experience two among time vector tracted and more vectors because the volume of acid or alkali added for the purpose of acidifying, or rendering alkaline, a solution is inadequate. Hence when rying or rendering anamon, a natural sa management of the seid or alkaline, the acid or alkali should be added whilst stirring with a glace rod until a drop of the solution removed on the end of the rod gives the required reaction with litmus paper. The method of preparing a perfectly Procedure for the identification of O.S.

[1] Determine the elements present by applying Tests 1-7 (pages (1) Determine the commons process of appaying access as under \$5.101. If only negative results are obtained assume O.S. contains C and II, and possibly O. Also apply Test 8, if applicable.

(2) Ascertain the class of O.S. by means of an appropriate acheme (see page 11), then follow the instructions given in order to identify O.S. as a particular member of the class.

Notes concerning the class schemes and sections.

(1) If a test depends on the production of a certain odour with which the student is not familiar, he should obtain some of the substance mentioned and compare its offer with that produced in the test, (2) Methods for the determination of m.p. and b p. are described

on pages 12-15. A refer apparates in Contract on page 17 to page acres in remos apparates on page 22 (Fig. 11). (3) It is probable that the majority of substances from for



# -SIS FOR ELEMENTS

facilion labes required for some of the tests may be prepared as follows:

Rest to a Banco dance with continued rotation, the centre of a Heat in a Binners issue with continuous relations the centre of a price of a off thin whited glass fabing, the control of districted districted and a control of the contro pice of soft into waited fines tuning 4-in outdoor connected and o in the fines is soft enough frames to take from the

long. When the glass is soft enough remove the tube from the control of a total length of 12 to -14 tn; allow to cool. tame according out to a total sength of 12 in-14 in., allow for cool.

Cut with a file in such a way that two wide tolers are obtained, Car with a his in such a way that two wide those are contained, as a such as a such a such a such a such as a such Marine 41 one end a lapering portion about f its long. Motate the there and in the titue unit sensed and continue heating and intelligential there is at the end of the table a remoded, rechoic

rotating until there is at the end of the libe a rounded, red-hot and the filter about it in deep. Remove from the fitting and the date and the liber. mass or feat about f in. deep. Hemore from the flame and blow. Dut that treetien of the smalless and exception while that the table is formed. dors the labe and a balb algoly where then the labe is formed.

All that period of the capitary take, which is of more or less

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than the capitary take, and the capitary take is the capitary take. All that Potting of the caputary tank, which is of more or twee Point determinations.

(1) structure a test for majorens.

Libert one end of about the of front copper wire into a core duel Abort one end of about 4 to of stout copper wave time a core and abort 4 to of stout copper wave time a core and the common of t bend about i in. of the other cod over so that it has practically particle of the wise. Heat the best cod of the wise. to the longer length of the war. Heat the heat end or the war. until it ceases to impart any colour to the many, then allow it to raise a little O.S. on a clean watch, then and dip the wire COL. There a little U.X. On a circum watch gives and dip the war follows of a cold a little Pile should be on the of the circum and the circu falls it: in the case of a saint a little pic about to on the ord of the wire in the lower course of or the the wire. Heat the end of the wire in the lower cuter edge of the mine in the lower cuter edge of the large. substance, has disappeared

ultitation has disappeared.

No firms reduce indicates the alleges of history (see note). No Direct colour indicates the absence of helicon (see mote).

A first colour is produced by taken in present, but the same of the colour is produced by taken in present, but the same of the colour is present to A firm colour is produced if ballyto is present, but the states

count is given by several substances which conclude no delivers on the several substances which conclude no delivers on the several se

ore The Erres colour cay be caused by a price extens become

J. Vo.s. to eliquid, to jet harry of h las 11, ed 3 e.e. de 7 y O. S. la Riguid, to fine lart of H is a 11, and 12 et al. 1. The state of the s

#### TESTS FOR ELEMENTS

If O.B. has not mixed with the water, add a drop of lodine so ion, shake, and allow to stand until the layers separate.

A violet-enfoured solution of locline in O.S indicates the above if O in the latter; if the colour of the solution is red or brown )B. may, or may not, contain O.

(3) Soda-lime test for N and Hg. (See notes.) If O.S. Is

(a) a solid, grind an amount of freshly ignited soda-lime which rould roughly fill the bulb of an ignition tube with about 1 of i sulk of O.S. Place sufficient of the mixture in the tube just t ill the bulb. Heat the bulb by rotating it over a small fiame an sold a piece of moistured red litmus paper about 1 in, from the mout of the tube.

If (i) the colour of the litmus paper is changed to a definite blu the presence of N in O.S. is indicated.

(ii) a grey sublimate is formed, rub it with the end of a match stalk. The formation of metallic globules indicates that O.S. contains Hg.

(b) a liquid, introduce into an ignition tube sufficient of it to ualf fill the bulb and add dry sods-lime to within about 1 in. of the nouth of the tube. Heat the tube, beginning at the top of the oda-lime layer and slowly proceeding downwards towards the bulb. Hold a piece of moistened red litmus paper about 1 in. from the nouth of the tube. If the colour of the litmus paper is changed o a definite blue the presence of N in O.S. is indicated.

JOYES.

(i) The absence of alkaline vapours on heating O.S. with sodaime does not indicate the absence of N in O.S.

(ii) If O.S. is a solid it should be noted if any odour (other than hat of O.S. or ammonia) is evolved during the heating with sodaime, as valuable information concerning the identity of O.S. may ften be gained in this way.

Thus an odour of

(a) phenol or a cresol indicates that O.S. is a phenolic scid or a alt or ester of a phenolic acid. If O.S. contains S a sulphonic cid, salt, or ester is indicated.

(6) an amine indicates that O.S. is a salt of an amine, an acyl erivative of a primary or secondary amine, an amino-acid, or a alt or ester of an amino-acid. If O.S. contains S an aminosulhonio acid or salt is indicated.

The lower aliphatic amines possess ammoniacal, fishy odours; the dour of pyridine is peculiar and unpleasant; and the commoner romatic amines porcess odours similar to that of aniline or of

(c) bitter almonds indicates that O.S. is a derivative of benzaldehyde (bisulphite compound, oxime, etc.), a mononitrobenzoie anid, salt, or ester (nitrobenzene evolved) or an amide (e.g. benzamide which yields benzonitrile).

(d) benzene or toluene indicates that O.S. is a simple aromatic

carboxylic acid, salt, or ester.

Whatever odour is detected the normal procedure for identification should be continued and tests shortened or omitted as far as the knowledge gained from the soda-lime test will permit.

(4) Alkali-sugar test for S and halogens and N in a limited number of cases. (Middleton, The Analyst, 1935, 60, 154.)

If O.S. is a solid, mix an amount equivalent to two or three times the bulk of a rice grain with about five times its bulk of alkali-sugar mixture. Introduce the whole into an ignition tube and add alkali-sugar

until, after tapping down, there is a column of the reagent about l in, long above the bulb.

If O.S. is a liquid, introduce two or three drops into the bulb of an ignition tube and add alkali-sugar until, after tapping down. the column of reasent extends to within about 4 in. of the mouth of the tube.

Hold the tube horizontally (by means of tongs) and heat it in a flame about 21 in. high, at first just beyond the column in order to prevent movement of the latter along the tube. Gradually extend the heating along the column, periodically turning the tube over to prevent undue bending. When a portion of the column equal in width to the flame is red-hot, hold the tube at an angle, so that while this portion still remains in the flame the heating is gradually extended until the bulb as well as the stem is in the flame. (It is essential that the whole column of reacent becomes red-hot before the organic substance in the bulb is heated.) Finally heat the whole tube to redness in a large fisme for a minute or more, then plunge it into 10 e.c. of distilled water contained in a dash. (If necessary, break up the tube by tapping it with the tongs.) Heat the contents of the dish to boiling, continue boiling with stirring for I min., then filter, (If the filtrate is not colourless repeat the whole process, finally beating the whole take more thoroughly.)

Apply the following tests to the alkalme filtrate:

(a) To I c.e. add one drop of an had acrease.

A brown or black prt. indicates that O.S. contains S. (Tracer a white pot.)

(b) To 2 c.e. add two or three drops of aq NaOII and one or two small crystals of FeSO, boil for I min, then cook Just acklify with cone. HCl, heat to boiling and cool. (If 8 has been detected, add one or two drops of an. FeCl, as the ferrio salt formed by atmospheric oxidation may have been reduced by the H.S liberated.)

A blue ppt, or colour indicates that O.S. contains N. (A green soln., if filtered, may leave a blue residue.) If a yellow or greenish yellow soln, is obtained apply Test 5

after Teets (c) and (d).

(c) To 2 c.c. add dil. H.SO, until the soln, is just said (if N or B is present, then dilute to 30 e.e. with water and boil down in a dish to 3-5 c.c.; cool), and add 1-2 drops of chlorine

water. If a yellow or brown soln, results, add about 1 c.c. of

chloroform and shake, -chloroform coloured brown indicates that O.S. contains Br.

violet If a violet chloroform layer is obtained, add CI water drop

by drop with shaking until the violet colour disappears. If the chloroform now has a brown colour, this indicates that O.S. also contains Br.

(d) To the remainder of the filtrate add dil. HNO, until the soln, is acid (if N. S. Br, or I, is present, add to the scidified soln, an equal volume of dil, HNO, dilute to 30 c.c. with distilled water and boil down in a dish to 3-5 a.c.; cool), then add 1 c.c. aq. AgNO,

A white curdy ppt. indicates that O.S. contains Cl. (Ignore a faint milkiness.)

Nore.

In the absence of sulphide, bromide and iodide, chloride may be detected in the presence of cyanide as follows: To the soln. (acidified with dil. HNO.) add I c.c. aq. AgNO, then add aq. mercurous nitrate (5%) with shaking until the black coloration first formed has completely disappeared.

If a white ppt, remains, this indicates that O.S. contains

Cl. (R. E. D. Clark, J.C.S., 1936, 1050.) (5) Alkali-zinc test for N. (Middleton, The Analyst, 1935,

60, 154.) (For advantages of this test see page 266.)

Follow the procedure described under (4) but using alkali-rine instead of alkali-sugar mixture.

me she atkaline filtrate apply only Test (b). The addition of in the alkaline



the presence in O.S. of As.

until the soln, is just sold. Pour I c.c. of the soln, into another t.t., arld 3 c.c. ammonium molybdate soln., shake, and allow to stand for 2-3 min. If (i) a yellow ppt, is formed, this indicates that O.S. contains P.

(ii) no yellow ppt. is obtained, heat to boiling and continue boiling for 1 min. or so. A yellow ppt, Indicates that O.S. contains As.

The following distinguishing test should also be applied: To the remainder of the acidified soln, add dil NH,OH until it is just alkaline. Pour the soln, into a dish and boil until a piece of red litmus paper momentarily immersed in the soln. is just no longer turned blue, keeping the volume between 3 c.c. and 5 c.c. by the addition, if necessary, of water. Cool the soln., and add 1 c.c. aq. AgNO, A pale yellow ppt, indicates P: a reddish ppt, indicates

Note A porcelain crucible is not advisable owing to the possibility of calcium phosphate being one of the constituents of the porcelain.

(8) Tests for halide, sulphate, and phosphate in solid compounds containing N.

If in addition to N (a) halogen has been detected, to 1-in. layer of O.S. in a t.t. add

5c.c. distilled water, heat to boiling and if O.S. has not dissolved continue boiling for | min. Cool and if any solid is present filter. To the soln, or filtrate add 2 c.c. dil, HNO, and then add I c.c. aq. AgNO. A white, pale yellow, or yellow ppt. indicates that O.S. is probably a halide salt of a base. (A similar result would be obtained with an acid halide of a

N-containing carboxylic acid.) (b) S has been detected, to 1-in. layer of O.S. in a t.t. add 5 cc. dil. HCL heat to boiling and if O.S. has not dissolved continue boiling for & min. Cool and if any solid is present, filter. To the soln, or filtrate add I c.c. ag. BaCl., A white ppt, indicates that O.S. is the sulphate of a base.

(c) P has been detected, to 3 c.c. NII, molybelate soin, add R.G. of O.S., shake and allow to stand for I min. - - stan of a mallow and fadinates at a OR is the

# INDEX OF CLASS SCHEMES

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## Note.

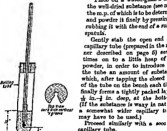
Augra.

In the heading on left-hand pages, those elements which may or
may not be present in the substance under investigation are in

## DETERMINATION OF PHYSICAL PROPERTIES

Determination of the melting point of a solid.

Fit up the apparatus shown in Fig. 1 and support the boiling tube by means of a clamp and stand so that the bottom is about 8 in above the bench. Into the boiling tube pour a suitable liquid (see note 1) to the depth of about 11 in.



Place on a watch-class a little of the well-dried substance (see note 2) the m.p. of which is to be determined, and powder it finely by pressing and rubbing it with the end of a suitable spatula.

Gently stab the open end of a capillary tube (prepared in the manner described on page 6) several times on to a little heap of the powder, in order to introduce the tube an amount of substa which, after tapping the closed of the tube on the bench each ti finally forms a tightly packed la d in. deep, at the bott

Proceed similarly with a seco capillary tube.

Remove from the boiling tube t cork and thermometer, and stro the moistened end of the latter alor

the lower portion of the capillary tube, then press the tube again the side of the thermometer so that the substance is opposi the middle of the bulb; the tube will adhere to the thermomete by capillary attraction.

Replace the cork and thermometer, gently heat the liquid wit mell flame, and note the temperature at which the solid change

to a transparent liquid (see note 3); an approximate ; m.p. is thus obtained.

Carry out a more accurate determination at once, by the procedure with the other capillary tube, raising, c degree, the temperature to which the liquid has fallen by organes, and temperature of many one organs was concerned as many fame (about \$ in, high) under the be the boiling tube until the m.p. of the solid is reached. the countries must sue to be to some the construction of the possible to reach, and not exceed or only to nucle practice is is possion to reach, and not exceed or only a exceed, a particular temperature. (With this form of app. execut, a paracular scapersoure. It is an ever or appearance the liquid is not stirred, the flame should not be kept timously under the boiling tube, or the thermometer will rea temperature which is actually below that of the liquid.) recuperature wames to accuming to destroy packed capillary tube and be used. NOTES.

(I) For temperatures up to about 210° medicinal parasita year at a suitable and reasonably as a liquid to use Jecus no successo and resonancy sais influent to use.

For temperatures up to about 200 fresh concentrated sulphuric for temperatures up to acoust you tress concentrated suppursual days be used, a crystal of KANO, being added to oxidise charred d may be used, a crystal of KAUs being added to oxidine charged the and prevent the sold from becoming discoloured. With

idea and prevent too seek more recoming associated. What ideal, however, there is risk of serious fujury should the tube liquid, however, there is that of serious injury shown the tube is, safer liquids are ceders of high b.p., s.g. butyl phthalate 3397.
It is essential that the substance be dry since even a trace isture may lower the m.p. considerably.

For methods of drying substances see page 19. for methods or drying submanics see page to.

(3) A pure compound metally has a sharp mp. i.e. it melas (3) A pure compound usually has a sharp mp. i.e. it melts make the make a sharp mp. i.e. it melts a sharp mp. i.e. it melts the make the m

npetery winin a ringe of about 1. Any impurities present the m p, and also render it indefinite, it the The strays more vise in the same are remore in minerance, e.e. in the form solid to liquid extends over a number of degrees. 

the arbeitance on heating undergo decomposition before the une studentes on assung undergo decomposation periore but is meshed, the decomposition products then acting as far. in reaction, the decomposition privates seen saling as mis-ses and forering the mp. Fren for these the method described see and lowering the in.p. Even for times the method described he introduction of the capillary take into the legid at a contract of the legic at a contract of the legid at a contract of the legic at a contract he introduction of the capuary time into the inquir as a state only a little below the mp. of the imbedancy enables Secure only a section activate cap, to co constance, same tos compound as exposed that temperature for only a short time before melting, and

thus only light decomposition occups a short time before mailing, and the constraint of the substance and the constraint of the substance and there thus only slight decomposition occurs. It too substance and sacra-momenter are placed in the odd liquid and the femperature raised story a much lower m.p. will be obtained. showly a much lower m.p. will be obtained.

[6] Since an ordinary thermometer may be inaccurate it should be made and the many thermometer may be inaccurate it should be made a range of pure substaines of known mp lan ...

## PHYSICAL PROCESSES

Filtration by suction.

Fold precipitates or crystale should always be filtered of from the legal present by exciton, since not only is the method much more rapid than ordinary filtration, but it also permits of the

almost complete removal of the limit.

(In the method described an inexpensive, ordinary type of conical funnel is employed. If available, a small porcelain Bichner funnel or a glass funnel with a plate of sintered glass, is more convenient.) Fit a filter flask (a thick walled conical flask with a short side tube for connection to a water pump) with a glass funnel (2 in. 2 in. diameter) by means of a lorsed rubber stopper. (Fig. 4.) Place in the funnel



a perforated porcelain disc. (1-in diameter is a convenient site.) Out a circle of dilter paper (see note, page 10) about virie the diameter of the disc, and fit it roughly in the funnel over the disc. Mostern the paper with water (or the solvent used for crystallisation), then press it flat with the fingers on to the disc, and mould the edges against the sides of the funnel. Connect the filter flash to the water pump by meas of pressure tubing, start the section, as pour the mixture of solid and liquid grade allow on the widdle of the filter page.

Pro. 4.—Pittes

Fleak.

### Note,

Smooth filter paper is preferable since the solid can easily be removed from it without being contaminated by paper fibre; also it withstands the pressure better than the coarse variety.

Washing of the fiftered solid in the funnel.

The filter flask should be disconnected from the pump whilst adding the liquid used for washing.

If the solid

- (a) is crude, use the liquid suggested for this purpose in the instructions given for the preparation of the compound, in order to remove the reagents employed.
- (b) has been crystallised, in order to remove mother liquor allow drops of the solvent, used for crystallisation, to fall over the

whole surface. CRYSTALLISATION the crystals are observed to dissolve readily, omit the crystals are observed to dissolve reachly, ourse either case connect the flask scan to the pump, a street case councer was mask again we see Pump, a successful minutes in order to remove as a tallisation of derivatives. unser the derivative from the filter funnel to a dr water two ucrystative from two inver numer to a or a day 100.c.o. conical flask, or if the amount is a dry t. Add sufficient of the recommended solvent corer the solid, and heat to boiling (see note 1). over the solid, and heat to bouling (see note 1).

If the solid does not dissolve completely, or almost com-If the solid does not dissolve completely, or almost companies of the amount of the solvent effer heating again, solution of the solution quite or almost complete. If any solid is present, filter hot (see note 3). Allow to cool, or if speed is essential, cool by olding the tube or flask in a stream of cold

ater, meanwhile shaking. If crystals do not partie, scrape the glass in contact with the aid with a glass tod; if still no crystale separate, or apporate of some of the solvent (see note I), and cool again. If the derivatire separates in the form of an oil whits the sold is warm, add a little more solvent, and beat again until a clear soln, is obtained, then How to cool spontaneously, stirring and stor to cool specialization, survivor and a glass rod from time to time. the off the expension and wash in the manney described on Page 16 as a sate manner described on page to Sass, and dry. (See "Drying of Substances." Page 19.1 Norm.

(1) Inflammable solvents, such as alcohol and actions should be beated for eraporated data service, known to neare for electrons, and packing the tube or flack in when,

toral by placing the tube or mast in water,
which has been raised to a suitable temperature, and from which
a manage of the first temperature, and from which the source of heat has been removed. A about de mes on some of heat has been removed.

In order to a world less of a sealthy volume solvent, substances, and a sealthy substances, and order to a rold loss of a reactly rotatile solvers, strong story should be bested with the solvers, strong one of the solvers as a flat fitted with a reflex condenser. (Fig. 5.) litter with a rettler condenser. (Fig. 5.)

(Fig. 5.)

\*\*Standard standly soluble in the solvent employed

(3) For the Elitation of a hot solm, a fined file

13 fannel, the stem of which has been cut off (Fig. 6), should be used;

also the funnel should be warmed, and the soln, kept as hot as possible during filtration. First, however, it is advisable to ave the soin, slightly and if any solid is readily deposited, to abl a hule more solvent and reheat, in order to prevent the adid crystallising out in the filter paper, the excess of solvent may be removed by evaporation (see teste 1) after filtration. When only a small amount of saled has been described it is rarely that any other procedure

will be necessary. To propers a flated filter fold a filter paper into four, balf open is and it wish each quadrant inwents on that the edge is parallel to the centre crosse A. Next double back each edge section, then

f I I such wate section backs relations the center, finally felt down the making erease as as to chitain a fin. Open out the fifter and fill oe h of the two pertengular flatings down the confin

Either of the following methods may be adopted:

(a) Use a suitable mixture of alcohol and water in the manner described for a single solvent.

(8) Dissolve the substance in the minimum quantity of fot alcohol, and then add water doop by drop with shaking until a turbidity just appears. Heat until a clear soln, is just obtained, and allow to cool alowly. If only a small proportion of the dissolved solid cystallase out, wheat the intrue until a clear soln, is again obtained, add more water and repeat the above procedure.

It an emulsion appears during the cooling, dissolve st by adding a few drops of alcohol and stirring, thus keeping the soln, clear until the temperature is sufficiently low for crystallisation to start. Once crystals have separated the mixture may be cooled and stirred in order to complete the

crystallisation rapidly.

(2) When a substance is moderately soluble in hot water, and readly soluble in cold alcohol, solution may be effected by boiling the solid with a volume of water insufficient for complete solution, allowing to cool somewhat, and then adding alcohol gradually with string or shaking until the remaining sold has dissolved. In this

way a large volume of solvent is avoided.

Glacial actic acid and water, or actions and alcohol may be employed in the same manner as alcohol and water.

Choice of a solvent.

If it is found necessary to recrystalize O.S. expression must be carried out in order to find a must be solvent. The commonest extent are written of actions and sicolob, duties more extent are written of actions and sicolob, features, perticular states. The procedure described under "Dystallization of deviratives" (page 17) should be followed, usua 2 µm, hayr of the powdered sold in a clean, day £5. A mustable solvent will be one of which 10 c.c. or less will describe the sold completely (or almost which 10 c.c. or less will describe the sold completely for almost advantage of the sold which will primit of the separation of the folk of the sold which will primit of the separation of the folk of the sold in crystallize form on coding and, if necessary, scraping the place to contact with the liquid with a place red.

Drying of substances.

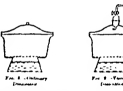
Lizaile.

Fromed as with an ethernal solu (are page 22).

Solute.

If it is present to dry a small preton of a detractes as finish as provide (or fix a mp, determination), proved as follows: Place a thin layer of the substance on a piece of smooth filter paper, and more the paper slowly from side to side in a borisontal plane, 2 in-3 in above a small flame. If the substance shows signs of melting, increase the distance between the paper and the fluor Feriodically move the substance about and turn it over with the blade of a penhnife. Continue the procedure until the ease with which the substance can be shaken from the knife blade fedicates that it is smith day.

Fairly rapid drying may be achieved, whilst other work is being carried out, by standing the watch-glass containing the mbelance, on the bench at a suitable distance from a triped on which a water bath, or a vessel on a gauze, is being heated by means of a Bunsen. If no vessel is being heated, a piece of asbestos board, or a grass with an asbestos-covered centre, should be placed on a triped on as to direct downwards the heat from a Bunsen placed broosth. A suitable position, i.e. one at which no melting, submarks, or discoloration occurs, may be found by trial with a small quantity of the adds.



In mose some the melanation may be suffly and regulty divid by quering face with given we a character block statisting lengthers? or so was for the house, as more to the Sains as practice. Public unity during two banking the mystacion should be turned over and supperform you as a points.

If your 3 with mention for up may be effected by planting the fact of the planting the fact of the send of the fact of the fac

those of benzene or petroleum ether) freshly cut pieces of paraffin wax are suitable.

A vacuum desicoator (Fig. 9) is much more efficient than the ordinary type, when exhausting it by missan of a water pump, a filter flask should always be fitted between the desicoator and the pump, in order to trap any sater which may rush back through the pressure tubing when the pump for any reason coases working. When opening the exhausted desicoator the art should be admissed slowly in order to avoid the dried substance being blown off the watch plans.

#### Extraction with ether.

Procedure for the extraction of an oil from a mixture with water, or of a substance from its solution in scater.

#### CAUTION.

In all operations where other is employed see that no flames are in the vicinity.

Cool the mixture or soln, to ordinary temperature, and pour it into a separating funnel (Fig. 10) 1100 c.c. capacity

will usually be suitable), then add to the mixture about 1 of its volume of ether (see note 1). Insert the stopper, and hold it in place with a finger of the hand which is holding the funnel. Invert the funnel and release the pressure by opening the tap. Close the tap, shake for about a minute, inverting the funnel and releasing the pressure from time to time. Allow to stand until two well-defined layers are formed (see note 2). Remove the stopper, and run off the lower layer (which may be required) through the tap into a braker. (The acueous solution drawn off is always saturated with ether, and should not be heated over a free flame.) Replace the stopper, and more the funnel about so that the ethernal soln. flows all over the inner surface. Allow to stand, then run



Fm. 10.--Separating Fannel.

off any lower layer which has collected (see note 3). Four the sterous shot, hough the peck of the famula into a dry 100 ca. wide-monthed fask, and connect the fatter to a water conferent, (Fig. 11). Immerse the lower part of the fask in het vater consisted in a beaker, Others the distillate in a class fask, and transfer it to a bettle for fatters one. (If the volume of the stransfer it to a bettle for fatters one. (If the volume of the stransfer it to a bettle for a fatter one.) Demove the fast its consistency of the fast fatter of other from the recoles by heating the fask, vationt the work, in belings water for a few minutes, and blowing air (by means of belings) over the resident.

#### North

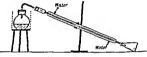
(1) If a large quantity of solution is to be extracted, add some Nat'l before the ether, ainer the latter is somewhat soluble in water but been so in brine , also many organic compounds are less soluble in brine than in water,

(2) If (a) the separation of the lavers is very imperfect, add

more other.

(b) a stable emulsion is formed, add a few drops of alcohol, which will usually disperse it.

(r) solul is present in the lower layer, add water; if this does not desolve it, filter the whole contents of the funnel, and pour the filtrate back into the washed-out funnel



Fro. II .- Distillation Amountus.

(c) Aqueous solutions: Heat the flask directly on a gause.
(b) Ether or other very volatile liquids: Heat the flask by partially immers in hot water contained in a beaker, and see that no flame are in the vicinity.

(d) the substance to be extracted is in solution in water extract several times with fresh quantities of ether and add together the several extracts.

(3) Washing and drying of the ethereal soln.

If it is necessary to wash the ethereal soln, free from acid, alkali etc., add about 3 c.c. of water for every 10 c.c. of ethereal soln., shake up, allow to stand, and run off the lower layer. Repeat this procedure until the wash water gives no test for the scid, etc., originally present. (The use of a large volume of water would soon greatly diminish the volume of the ethereal layer, since ether is somewhat soluble in water.)

When it is necessary to determine the b.p. or m.p. of the extracted compound, the ethercal soln, should be dried before distillation, since ether dissolves a little water. The drying agent to be used will depend on the nature of the compound dissolved in

the ether.

For most substances fused CaCl, may be used; this substance, however, forms addition products with alcohols and bases, hence

anhydrous K<sub>2</sub>CO<sub>2</sub>, or anhydrous Na<sub>2</sub>SO<sub>4</sub>, is employed for these classes of compounds.

classes of compounds.
Add a Bittle of the drying agent to the ethereal soin., and allow
to stand. If the drying agent sbutchs sufficient water to dissolve
it, run off the sq. soin, and and more of the drying agent. Before
distilling, pour off the ethereal soin, from the solid drying agent.
Ad dry ethereal soin, should never be freely exposed to the ans,
as the evaporation of the ether causes a considerable deposit of
moisture on the soin.)

these colours is obtained, apply Test (b). (b) To 5 drops of O.S. in a t.t. add 3 c.c. of 10% sq. KOH (NaOH is not suitable as it forms a sparingly soluble sodium salt with some compounds). Close the month of the tube and shake vigorously for 1 min. If any O.S. has remained undissolved add 3 c.c. of water, then shake and filter the contents of the t.t. unless they consist of a gelatinous mass with a lather (suggesting that O.S. is oleio acid) in which case apply Test 3. Acidify the soln. or filtrate with conc. HCl. If a white emulsion or an oil is obtained, indicating that O.S. is either a phenol or a carboxylic acid, apply Test (c); if no such result is obtained, apply Test 2.

(c) To 1-in. layer of solid NaHCO, in a t.t. add 5 drops of O.S. and 5 c.c. of cold water. Close the mouth of the t.t. and shake vigorously for 1 min., allowing any evolved gas to escape. Carefully acidify the soln. (previously filtered if any

O.S. had remained undissolved) with cone. HCl.

If (i) a white emulsion or an oil is produced, this indicates that O.S. is a carboxylio acid or contains a carboxylic acid, probably formed by the oxidation of an aliehyde or by the hydrolysis of an ester. Apply Test 2.

(ii) no white emulsion or oil is obtained, determine the b.p. of OS. and refer to the list of b.p s of monohydrle phenols (page 46).

(2) Tests for aklehydes, polymers of aklehydes, acetals, and betones. Follow the appropriate procedure according to whether OS is miscible with water (below) or not miscible with water (page 27).

#### O.S. miscible with water.

(a) To 2 c.o. of the cold prepared an mile, of OB, add 2 co. of Schiff's reagent, shake to mix and allow to stand for 2 min. Do not heat the mixture.

If (i) a deep walet or deep sed calour is obtained, one under A (page 54).

(a) no privat or only a funt sal colour is preduced, then to 2 ca. of the prepared on with of till will me drop of arress 2% Hr3 brand the \$1 to belling water for 2 mis. (see mide below), then jumies it still eral tre statesta. Att 2 a c of the bill a magnit and thake to mix. If within 2 min. a deep violet or deep red colour is obtained, see under A (page 54); if no colour or only a faint red colour is produced apply Test (6).

#### Nors.

The formation of a white emulsion, changing in colour through yellow and orange to red brown, auggests that O.S. is furfuryl alcohol and Test 4 should be applied.

(b) Add 2 drops of O.S. to 2 o.e. of \$\frac{1}{2}\%, aq. soln. of sections nitroprusside, then add approx. 2N. NaOH, drop by drop, until the soin. when tested with red limus paper gives an alkalme reaction. (If a red colour is produced with one drop of NaOH no further addition of alkal is necessary.)

If s wine-red or orange-red colour (which soon changes to yellow) develops in a few seconds, see under B (page 55); if only a yellow colour develops, apply Test 3.

#### O.S. not miscible with water,

To j.in. layer of 2:4-dinitrophenyhydradne in a dry t.t. add 2e.c. of alcohol, then add j.c. of conc. HSO. Warm and shake in order to dissolve all the solid. To the warm soln, add e.c. of O.S. and allow to stand for 5 min. unless a ppt, forms in a shorter period of time. (The warm HSO, will depolyments as a shorter period of time. (The warm HSO, will depolyment allowed the polyment of the dischapte will the respect.) Finally cool and ababe hydrowing the mean control of the control of the discharge with the restain it and proceed as indicated under "O.S. not miscible with water" (page 57); if there is no ppt, apply Test 3.

To 2 c.c. of alcohol add, using a dropping tube, one drop of O.S. and shake round, then add 1 or 2 drops of phenolphthalein soln. From a similar dropping tube add approx. NaOH, drop by drop,

shaking after each addition until a red colour, which persists for 1-2 sec., is obtained, or until 10 drops of alkali have been added. If after the addition of

(a) 10 drops of sikali the soln. is colourless, see "Carboxylic acids, etc." (page 67).

(b) 10 or force drops of alkali a red colour, persisting for 1-2 acc, was obtained them, if the soil, is still red, heat it to boiling and notice if the colour disappears or not. The disappearance of the red colour, either on standing or on boiling the soils, suggests that O.S. is an ester a hydroxyl group may also be present in the molecule. C, H, [O]

(4) Test for akohole. See caution below.

To I c.e. of acetyl chloride in a dry t.t. add about i c.c. of OS

If there is no immediate reaction, watch for 2 min.

If (a) a vigorous reaction occurs (i.e. bubbles are freely evolved) with evolution of HCl fumes, see "Alcohols" (page 33).

(b) there is no reaction, see "Esters, ethers, and hydrocarbona" (page 99).

CAUTION.

Alcohols, which are miscible with water, react immediately with acetyl chloride, often with explosive violence, and the mixture may be expelled from the tube.

NOTES.

(1) Since a reaction with acetyl chloride might be due to the presence of water in O.S. it is advisable to allow a quantity of O.S. to stand in contact with about | of its volume of freshly ignited K.CO. for 1 hr. or more and then to repeat the test with acetyl chloride. In practice, however, it will rarely be found that members of the classes of compounds not yet detected (i.e. esters of non-hydroxy acids, ethers, and hydrocarbons) will contain sufficient water to give any appreciable reaction with acetyl chloride.

(2) With furfuryl alcohol, in addition to the vigorous reaction, the mixture becomes violet, then very dark green.

O.S. solid.

If O.S. contains a metal proceed as indicated under "Metal present " (page 31), otherwise follow the procedure below.

No metal present.

Tr o.s. (a) possesses a pronounced yellow, orange, red, or green colour,

see "Coloured solid compounds" (page 95).

(b) is an amorphous white powder, add R.G. of it to 5 c.c. of water in a t.t., heat to boiling, cool, and nearly fill the t.t. with water. Add one drop of iodine soln., close the

mouth of the t.t. and mix the contents by inverting. If a deep blue colour is obtained see "Starch " (page 93), or if there is no blue colour prepare an aq. soln, of O.S.

in the manner described below and apply the class tests in the order given.

(c) does not possess the properties described under (a) or (b). prepare an aq. soln. of it in the manner described below and apply the class tests in the order given.

Preparation of an ag. soln. of O.S .. To a measured 1 in layer of powdered O.S. in a dry 1 in. i.t. add 5 c.c. of distilled water and heat to boiling. If O.S. has s dissolved completely continue bolling with shaking for † min. et the contents of the st. A collection of the state of the stat

### CLASS TESTS

ests for phenolic compounds and carboxylic acids. .

To 2 c.c. of the cold prepared sq. solz. of O.S. add one drop

of eq. FeCl,

If a blue, violet, red, or deep green colour is observed, see "Phenolic compounds" (page 44); if none of these colours is obtained proceed as under (i) if O.S. is readily soluble in

is obtained proceed as under (i) if O.S. is readily soluble in water or as under (ii) if O.S. is sparingly soluble in water.
(i) Add R.G. of O.S. to 2 c.c. of dil. NH OH and shake

for 1 min. If an immediate yellow colour, which changes to radbown; is observed, determine the mp. of 0.8, and see "Folynydro phenola" (page 49); if no such colours are produced, then to a solution, obtained by boiling R.G. of 0.8, with 2 c. c. of distilled water

and cooling, add 5 drops of approx. N NaOH, shake

and then add a drop or two of phenolphthalein soln.

If the soln is colourless, proceed as indicated under 
"Carborylic acids" A (page 70) or if a red colour is

"Outdoord about Test 2 a.

(ii) To 1-in. layer of O.S. in a t.t. add 2 c.c. of alcohol, heat just to boiling, then cool and add 1-3 drops of eq. FcO. If a blue, violet, red, or green colour is observed, see "Phenolic compounds" (page 41); if

nono of these colours is obtained, apply Test (b).

Jein Jayer of prowdered O.S. in a st. add 3 a. o. of approx.

K. NOH, closs the mouth of the tube and shake vigorously

I min. If any O.S. has remained undissolved add 3 a.c.
water, then abake and filter the contents of the t.i.
slift the soln or filtrate with cone. HCl. If there is no
meduate ppt., apply Test (c); if an immediate ppt. is
not the topic apply Test (c); if an immediate ppt. is
tained, inducting that O.S. is when a phenol or a carboxylie

J, then to j-in. layer of powdered O.S. in a t.l. add an
all bolk of bold NaHOO, and 6 c.a. of cold water. Close
mouth of the tube and shake vigorously for I min.
wing any criving gas to exappe. Filter if any solid is
sent and carefully acidify the soln. or filtrate with cone.

If a ppt. is obtained see "Cubroxylie acids" under B.

C. H. [0]

30

(page 74); if there is no ppt. determine the m.p. of O.S. and refer to the list of m.p s of monohydric phenols (page 47).

(c) Add R.G. of O.S. to 2 c.c. of alcohol heat just to boiling. then cool. Add 5 drops of approx. NaOH, shake and then add I or 2 drops of phenolphthalein soln. If the soln. is colourless, determine the m.p. of O.S. and refer to the list of m.p.s of carboxylic acids (page 76) or if a red colour is produced apply Test 2 b. (This test (c) is for the detection

of higher fatty acids which would not have been indicated in the tests under (b).) (2) Tests for aldehydes, polymers of aldehydes, and ketones.

(a) To 2 c.c. of the cold prepared aq. soln. of O.S. add 2 c.c. of Schiff's reagent, shake to mix and allow to stand for 2 mix. Do not heat the mixture. If a deep violet-red colour is produced, see under F (page 64); if no colour or only s faint red colour is obtained, apply Test 3. (b) To 1 in. layer of O.S. in a t.t. add 5 c.c. of dil. HCL Heat to boiling, continue boiling with shaking for 15 sec., then

cool; filter if any solid is present. Add the soln, or filtrate to a soln. obtained by boiling R.G. of 2:4-dinitrophe hydrazine with 2 c.c. of dil. HCl and cooling. If a pp obtained, see under "Solids" (page 62) or if there is ppt, proceed as under (c). just to boiling, then cool and shake. If no solid is prest use this soln. for the test given below: if solid is prest add another 3 c.c. of alcohol, heat to boiling with shaki then cool and shake. Filter if any solid is present and t the soln. for the following test :- To 1-in. layer of 2:

(c) To 1-in. layer of O.S. in a dry t.t. add 2 c.c. of alcohol, I dinitrophenylhydrazine in a dry t.t. add 21 c.c. of slooh then add I c.o. of conc. H.SO. Warm and shake in ord to dissolve all the solid. Add the prepared alcoholic sol of O.S. and if no ppt, is formed heat just to boiling at allow to stand for 5 min. Should no ppt. form, cool, an acrape the inside of the tube in contact with the liquid wit a glass rod for a minute or so, then allow to stand for further 5 min. If a yellow, crange, or red ppt, is obtained proceed as under "Solids" (page 62), or if there is no ppt apply Test 4.

(3) Tests for certain glucosides and carbohydrates, readily soluble in water. In a dry t.t. place R.G. of O.S. and add one drop of cone, H.50.

If a crimson colour is produced see "Glucosides" (page 94); if

#### CLASS DE'

no such colour is obtained app

as follows:—Dubte the prepared as, solo. of O.S. with volume of water. To 2 c. of this dubted solo. add 2 Molisch's referred. (1972 point, of a naphthol in sloobol) as Carefully pour the following of a naphthol in sloobol) as down the side of the tube and slow to stand for 2 mi red-violet ring is obtained where the two layers meet and or the whole mitture becomes violet-red and a dull blue-vi forms, see "Carbohydrates" I (page 91); if no such obtained see "Soldt sloobols" (pages 38, 43)

(4) Test for certain esters, lactones, and peroxides, sparingly in water

To j.in. layer of O S. in a t.t. add 3 c.c. of 20% aq. Kt to boiling and continue boiling with shaking for 1 min.

If (a) a deep brown or green colour is obtained, suggest O.S. is an ester of a polyhydric phenol, proceed as it under C (was 109).

(b) the liquid becomes yellow or remains eclourless, if any solid is present thicke with an equal v water and filter. Acidify the soli. or filtrate w RCI; cool and shake. If there is no ppt, see eithers, and hydroactbona "(page 69); if a ppt, is this suggests that O.S. is a lactone, peronde, or a sparmply soluble acid, hence see "Lactones coides" (page 60); or if O.S. is not one of the cothere listed proceed as indicated under "Exteroxyle acids," "O.S. solid "(page 107).

#### Metal present.

Procedure :-

(a) To 1-in. layer of O.S. in a t.t. add 2 a.e. of dd. I effervescence of CO<sub>2</sub> indicates that O.S. is a metal onate. If there is no effervescence, apply Test (b)

(b) To ½-in. layer of O.S. in a t.t. add 5 c.o. of distill and heat to boiling with shaking. Test the soln. or with both blue and red litmus paper.

If an acid, neutral, or slightly alkaline reaction is see "Salts of carboxylic acids" (page 87). A alkaline reaction will usually indicate an alkonide (see or a phenoxide (see page 53).

## ALCOHOLS

These are detected by the rigorous reaction, with evolution of HCl furnes, on adding to acetyl chloride. (Test 4. Scheme I.) Procedure for Liquid alcohols:—

If O.S. is

(a) completely miscible with twice its volume of water, pour I c.c. of O.S. into a dry t.t., add an equal volume of ether, close the mouth of the tube with the thumb and invert twice. If O.S. is completely miscible with the ether, see "Mono

hydric Alcohols" (below) otherwise see "Polyhydri Alcohols" (page 41), not completely miscible with twice its volume of water, see "Monohydric Alcohols" (below).

# (b) not completely miscible with twice its volume of water, see "Monohydric Alcohols" (below). MONOHYDRIC ALCOHOLS

....

Liquid alcohols.

Procedure for the identification of O.S.— Determine the h.p., then refer, in the appropriate sub-section according to whether O.S. is completely miscible with twice it values of water [1998 33], floats on water [1998 35], or sinks in water [1998 37] to the list of the, so of alcohold, if one of the hya is identical with, or ness to, that of O.S., apply the test circum for that alcohol.

In many cases sufficient proof of identity will be obtained by oxidizing O.S. and identifying the oxidation product; in other cases the identity of O.S. may be confirmed by the preparation (see page 267) and determination of the m.p. of a derivative.

If the properties of O.S. are not identical with those of one of the alcohola in the lists, proceed as described under "Esters of carboxylio acids" (page 100), as O.S. may be an ester of a hydrox acid not included in the lists.

O.S. completely miscible with twice its volume of water.

B n.

65° Methyl alcohol. CH OH.

(a) Pour 2 drops of O.S. into a dry t.t. Bend abor 2 in. of stout copper wire into a compact form a

C.

(b) Carry out the exidation test (page 28).
78° Ethyl alcohol. CH, CH, OH.

82° (so-Propyl alcohol. (CH<sub>3</sub>),CH OII.

tert-Butyl alcohol, in.p. 25°. (CH<sub>2</sub>),COH.)
(a) To 1 c.c. of O.S. add twice the volume of 20% aq.
KOH.

Complete miscibility with the aq. KOH indicates that O S. is ethyl alcohol. Carry out the oxidation text (page 33). If O S. floats on the aq. KOH, apply Text (b)

- (b) To 2 c.c. iodine soln, add one drop of OS, then add aq. NaOli, drop by drop, until the deep brown colour changes to pale yellow. (2-3 dreps of aq. NaOli will normally be required.) The microliate formation of a pale yellow, finely divided ppt. of iodoform, with characteristic edox, indicates that OS. is ico-propyl alcohol. Carry out the oxidation test (page 38).
- (c) To 1 o.o. of 0.S. add 4 c.c. cono. HCl and shake.

  The formation of an insoluble liquid chloride, which on standing separates as an upper layer,
- indicates that O S. is tert butyl alcohol.

  96 Allyl alcohol. CH, : CH-CH, OH. Very pungent odour, resembling oil of mustard, irritating action on the eyes.
  - (a) Add 2 drops of O.S. to 5 c.c. Br water, —instant removal of brown colour, due to the presence of a double bond in the molecule of O.S.
  - (b) To 1 c.c. of O.S. add 5 c.c. dichromate mixture, —instant reduction, acrylic aldehyde (acrolein) formed with a still more irritating odour.

97° n-Propyl alcohol. CH .- CH

Carry out the oxidation test (page 33).

124° Ethylene glycol mono-methyl ether. (Methyl Cellosolve)
CII, OH

CH O Me

135° Ethylene giycol mono-eth

(page 41)

.

Methyl lactate. CH, CH(OH) COO Me) 140 Ethyl lactate CH, CHOHI COO Eth

of norms. University of the st. add 2 on come. His Oo warm cautionaly with shaking until the

neoth reasonably were assumed as well is pale yellow, then cool. And 2 drops of a 59 alcoholic solz, of gusison, in eye securious som, or guaracte, intense red colour, indicating a lactate.

(3) To identify the alkyl radical proceed as described 170° Furfuryl alcohol. CH : CH 10 Meeterly ton success face and a mercan to meeter the first of carboxytic acids " (page 190). CH.OH

(a) Add 1 drop of O.S. to 2 c.c. dil HCl. then seatly

warm and ename,
-white standard, changing to yellow, then crange, white equipon, changing to years, then crange, finally to red-brown. Dark coloured oil separates.

(Due to regulate tion of the alcohol by the distressed of sections at the alcohol by the distressed of the alcohol by the distressed.) (b) Boil 1-th layer of O.S. to a t.c. and hold a wooder tout Pur. myet or v.c. on a set pour most a western that first the various.

matchatalk turned binish green Ethylene glycol mozo-scetate. CH: OH

(a) Dissolve I a.e. of O.S. in 5 c.e. dil. MCI, pour the solar into a percelain dist, beat to belling and contains bolling for I min. Add dil MH OH until

the sin is alkaline, boil until neutral, then cool. To 2 cc. of the soln, add an equal return of aq. FoCl.

who.rd colour (riewed through the depth of the liquid) due to the formation of MI sociate (b) (kery out Tests (a) and (b) under "Physics Strates

(1950 41), when similar results will be obtained (c) Percentage acrif radical = 413. (For method of

Preryl diacetate (Diacetin), CH-OAe Paint acrid

odour. Somewhat w

- (a) Carry out Tests (a) and (b) under " Glyceryl monoacctate" (page 42), when similar results will be obtainet
- (b) Percentage acetyl radical = 48.8. (For method of determination ace page 43.1

O.S. floats on water.

B.D.

100° sec-Butyl alcohol. CH, CH, CH-OH

CH, CH-OH

108° iso-Butyl alcohol. (CH, cH-CH, OH)

(a) To 2 c.c. iodine soln, add one drop of O.S., then add aq. NaOII, drop by drop, until the deep brown colour changes to pale yellow. (2-3 drops of aq. NaOII will normally be required). Allow to stand for a period of time not exceeding 2 min. The formation (usually in about I min.) of a pale yellow, finely divided ppt. of iodoform, with characteristic odour, indicates that O.S. is

sec-butyl alcohol.

(b) Carry out the oxidation test (page 38). 117º n-Butyl alcohol. CH, CH, CH, CH, OH.

Carry out the oxidation test (page 38). 131° iso-Amyl alcohol. (CH.), CH-CH. CH.OH. Disagrecable odour : provokes coughing.

Carry out the oxidation test (page 38).

Nors. The amyl alcohol of commerce, obtained from fus oil, is a mixture of iso-butyl carbinol, b.p. 131°, an sec-butyl carbinol (active amyl alcohol) b.p. 129 both of which are primary alcohols.

138° n-Amyl alcohol. CH, [CII, 1, CH,OH, Carry out the oxidation test (page 38).

160° cyclo-Hoxanol. (Hexalin.) си — си си оп CH

Oxidise to adipie acid, m.p. 150° in the manner described under "cuclo-Hexanone" (page 60). 188° m. Butyl lactate. CH4-CH(OH) COOC.11.

Proceed as described under " Methyl lactate " (a) and

(b) (page 35). 197º Linslol. Cull .. Oil. Pleasant colour.

218° Terpineol. Cull 17 OH. Olour of lime.
Caronellol. Cull 11 OH. Olour of lemon rind. "Odour of rose and geranium. - A ---tal P

Add one drop of O.S. to 1-in. layer in a t.t. of a soln.

of Br in CCl.

-immediate decologisation.

-immediate decolourisation.

Pour the contents slowly out of the tube,

-no copious evolution of HBr fumes, indicating that addition, and not substitution, has occurred.

O.S. sinks in water.

B.p.

205\* Benzyl alcohol. C<sub>4</sub>H<sub>4</sub>CH<sub>4</sub>OH. Faint aromatic odour. (Unless freshly distilled possesses a slight bitter simond odour, due to the presence of benzaldehyde, formed owing to oxidation by the nin!.

(a) To 1 c.c. of O.S. in a t.t. add 1 c.c conc. HCl, shake, then immerse the end of the t.t. in boiling water.

-mixture becomes clear, and in about i min, a white emulsion suddenly appears, due to the formation of bensyl chloride. On standing the benzyl chloride separates as a colourless upper layer.

(b) To 2 cc. dil. HNO, (1 HNO, 4 H<sub>2</sub>O) in a t.t. add one drop of O.S. and stand the t.t. in boiling water for 2 min.

-pale yellow emulsion, and strong bitter almond odour of benzaldehyde.

(c) Oxidation to benzoic acid, m.p. 121°.

In a 100-ca. flask place the equivalent of a ½-in layer in a t.t. of sold KMnO<sub>2</sub> and 30 cc. water. Heat to boiling, allow to go just off the boil, then add 1 cs. of O S. Allow to stand, shaking round periodically, until the purple colour has disappeared, root. Filter and actify the fiftest with the colour of the colour of

220° Phenylethyl sloobol. C,H,CH,CH,OH,Oh. Odour of roses, (a) Gives no emulsion in 1 min. when treated as described shore under "Benryl sloobol" (a). On removing the tt. from the water and allowing to stand an upper layer (usually nink) is formed.

(b) The aromatic nature is shown by the formation of benzoic acid when oxidised in the manner described under "Cinnamyl alcohol" (b) (page 38).

n

11

Readily unfable in water.

97. are there almost collins out

To I se of autem OS in a st will 4 oc. come. HO. and shake-appear herer of perbutel chierile 4000404

Involuble to water

ita

23' Clansmyl almhot. Call, CH; CH CH, OH, bp. 250'.

Herwinth Like reloug. (a) To i.in. layer in a t t. of a soln, of Br in CCl, add

one drop of O.S. and shake, -deep brown colour dissprears almost instantly owing to the presence of a double bond in the molecule of O.S. (3) The aromatic nature is shown by the axidation to

benzoie acid, m p. 121°, as follows :- In a 100 c.c. flack place the equivalent of a 1-in layer in a tt of solid KMnO, and 50 c.c. water. Heat to boiling. remove the flame and add gradually I c.c. of molten O.S., shaking round after each addition. Allow to stand, shaking round periodically, until the purple colour has practically disappeared. Cool and pass in SO, until any purple colour and the brown ppt. have disappeared. Heat to boiling, filter to remove any oil present, then cool. Filter off the solid, wash it with cold water, dry, and determine the m.n.

#### OXIDATION TEST FOR ALCOHOLS

Identification of primary and secondary alcohols by recognition of the aldehyde, acid, or ketone formed by exidation. Procedure (for alcohols which are completely miscible with twice

their volume of water, or which float on water) :-

Fit up an apparatus consisting of a corked 50 c.o. distilling flask connected to a water condenser.

Disconnect the flask and pour into it 1 c.c. of O.S., add two or three pieces of porous pot, then pour in 20 c.c. of dichromate mixture. Quickly cork the flask and connect it again to the condenser. Use a t.t. (marked to indicate the space occupied by 7 c.c.) as the receiver. Distil until 7 c.c. of liquid is present in the receiver, noting when 2-3 c.c. is present if there is a layer of oil on the surface of the distillate.

If O.S.

- (a) is completely miscible with twice its volume of water, apply Schiff's test under A.
  - (5) floats on water, proceed as under B (page 40).

#### A. Note.

A thin layer of oil on the surface of the distillate indicates that O.S. is n-propyl alcohol. The distillate, however, should be shaken up and the following test applied.

Schiff's lest.

To 1 c.c. of distillate add an equal volume of Schiff's reagent, shake to mix and allow to stand for a period of time not exceeding 2 min. (The mixture must not be heated).

- If (1) a deep riolet-red colour is obtained (indicating that an aldehyde has been formed and therefore O.S. is a primary alcohol) proceed with the tests under "Akiehyde present."
  - (2) no colour, or only a faint pink colour is obtained, apply
    the following sea for formic acid ~ 20 1-2 cc. of distillation
    following sea for formic acid ~ 20 1-2 cc. of distillation
    following sea for formic acid ~ 20 1-2 cc. of distillation
    following sea for the following sea for following
    following sea for formic acid following sea formic
    following sea for formic acid following
    following sea formic following sea formic
    following sea formic following
    following sea formic following
    following sea formic following
    following sea formic following
    follow

A white ppt. of Hg,Cl, indicates that formic acid is present in the distillate and therefore O.S. is methyl slochol. If no ppt, is obtained proceed with the tests

under "Aldehydo absent " (page 40).

#### Aldehyde present.

To 2 c.c. of distillate add an equal volume of 20% aq. KOH, abake and allow to stard for I mis. Note if the mixture remains clear, or if a white emulsion is formed; in either case heat to boiling and continue boiling for j min.

If there is obtained

(a) in the cold a clear soln, and on bosing a reliew ppt, which changes to crange, with a desgreeable colour, the prepence of acetaldehyde in the dutillate is indicated.

of acetalicayde in the distillate is indicated.

Apply the following confirmatory test for acetaldehyde:—
To 2 e.c. of distillate add an equal volume of 1% aq.

modium nitroprusside, then add 1 e.c. aq. NaOH.

—wine red colour.

The presence of acetaldebyde in the distillate indicates that

O.S. is ethyl alcohol.

(b) in the cold a white emulsion, and on boiling a yellow colour

C. H. [0] (no orange ppt.) and a disagreeable odour, the presence of propaldehyde in the distillate is indicated. On adding to 2 c.c. of distillate an equal volume of 1% aq. sodium nitroprusside, then 1 c.c. aq. NaOH, only an orange yellow colour will be obtained. (Difference from acetaldehyde which gives a wine-red colour.) The presence of propaldehyde in the

distillate indicates that O.S. is n-propyl alcohol. (c) a clear, colourless soln. in the cold and after boiling, apply the following test for formaldehyde :-To 2 c.c. of distillate add R.G. of resorcinol, then pour

2 c.c. conc. H,SO, (from another t.t.) carefully down the aide of the tube.

-red ring at junction of liquids, white ppt. (which changes to violet red) forms in the aq. soln.

The presence of formaldehyde in the distillate indicates that O.S. is methyl alcohol. The test for formic acid described under 2 (page 39) should also be applied.

40

Aldehude absent To 2 c.c. of distillate add an equal volume of 1% aq sodium nitroprusside, then add 2 drops aq. NaOH. If a wine-red colour develops, quickly acidify with acetic acid. (If the mixture is not

acidified, the red colour will change to yellow.) A wine-red colour indicates that a ketone is present in the distillate, hence O.S. is a secondary alcohol.

Wine-red colour, turned violet-red by acetic acid indicates that acetone is present in the distillate, hence O.S. is iso-propyl alcohol.

Wine-red colour, little affected by acetic acid, indicates that methyl ethyl ketone is present in the distillate, hence O.S. is sec-butyl alcohol,

For further distinction, repeat the exidation and prepare a 2:4-dinitrophenylhydrazone, using the whole of the distillate. (For preparation see page 66)

m p. 126° 2:4-Unitrophenylhydrazone of acctone " methyl ethyl ketone, m p. 111° \*\*

NOTE. asec-Butyl alcohol (floats on water) is included here in order

to render the scheme suitable for the detection of an alcohol in ag, soln , e g the dutilists obtained from the hydrolysis products B. A layer of oil will usually to present on the surface of the

distillate. Shake up the distillate and apply Schiff's test. (See A deep violet-red colour indicates that the oil is a sparingly soluble allelytide, bence O.S. is a primary alcohol. If no colour, or only a faint colour is obtained, proceed as indicated under "Aldelytic absent" (page 40). The presence of methyl eithyl ketons in the distillate indicates that O.S. is accounted alcohol.

POLYHYDRIC ALCOHOLS

#### POLYHYDRIC ALCOHOLS

O.S. liquid.

Procedure for the identification of O.S.

A determination of b.p. will not be usually worth while, owing to decomposition of O.S. at high temperature, or to the presence of water, hence apply the following test:—

To i.in, layer of O.S. in a dry t.t. add KHSO, crystals to a depth of i in.; heat.

If there is obtained

(a) a pungent-smelling, highly irritating vapour (acrolein) see
 "Glycerol" (page 42).
 (b) a strong odour of actic send, followed on further heating

by a pungent-ameling, highly irritating vapour (acrolein) see "Glyceryl mono-acetato" (page 42).

(c) no puncent odour, see "Ethylene glycol" and "Diethylene glycol."

Bp. CH,OH

197\* Ethjiene glycol. CH,OH

CH.-CH,-CH,OH

215° Diethyleno glycol. O'CH, CH, OH CH, CH, OH (a) Dissolve I-in, layer of O.S. in a 4.4, in 10 c.s. aq.

Na<sub>2</sub>(O<sub>2</sub>. Four the sola, into a loding table in which has been placed the equivatest of a j.in, layer in a 1.1 of sold KMnO<sub>2</sub>. Bud for 1.2 min, then filter, (Shoull the Linux be purple in colour, desirent the stress of KMnO<sub>2</sub> by the adution of 11<sub>1</sub>(O<sub>2</sub>). Anally 2 is not the first switch the colour of the stress of the first switch of the colour of the colour of the colour of 3 drays of an Call. —white pr. 4 of Oa craise.

(i) Into a dry t t. poor 5 drops each of O.S. and come. 11,50, and heat until the mixture is a moderately

deep brown colour. Cool, carefully dilute to 5 c.c. B.p. with water, then add solid NaOH with shaking until the mixture is alkaline; boil, -characteristic, disagreeable odour of aldehyde regin. (c) Preparation of a benzoul derivative. Into a 100-c.c. flask pour 1 c.c. of O.S., 10 c.c. of acetone and then 5 c.c. of benzoyl chloride. Add 50 c.c. aq. NaOH (the first 10 c.c. or so

C. H. [01

gradually with cooling and shaking, then the remainder all at once); cool. Cork the flack and shake until the odour of the benzoyl chloride has practically disappeared: Filter off any solid present, wash it well with cold water, crystallise from

alcohol, dry, and determine the m.p. Ethylene glycol dibenzoate, m.p. 73°.

Diethylene glycol dibenzoate does not solidify.

decomp. Glyceryl mono-acetate (Monacetin). CH; O Ac

сн-он

CH. OH

Somewhat viscous.

42

(a) See the test under "Procedure for the identifica-

tion of O.B.," result (b), page 41.

(b) Dissolve 1 c.c. of OS. in 5 c.c. dil. HCl, pour the soln, into a porcelain dish and boil for 1 min. Add dil. NH OH until the soln, is alkaline, boil

until neutral, then cool. To 2 c.c. of the soln. add an equal volume of aq. FeClas -wine red colour (viewed through the depth of the liquid) due to the formation of NII4 acctate. (c) Percentage of acetyl radical = 32.1. (For method

of determination see page 43.)

290° Glycerol, CH OH Extremely viscous,

cn-on

CII. OII (4) See the test under " Procedure for the Mentifica-

tion of OS.," result (a), page 41. (b) Dimolre i-in. layer of boras in a st. in 20 e.s. water. To 5 ee. of this win. will a drop of

phenolphthalds sols , then ac .....

—red colour disappears; reappears on warming and disappears again on cooling.

(c) Prepare the benzoyl derivative in the manner described under "Ethylene glycol" (c), page 42.

O.S. solla

M.p. 166° d-Mannitol. CH,OH [CH(OH)], CH,OH. Readily soluble

d-Mannitol. CH4OH [CH(OH)]4 CH4OH. Readily soluble in water; insoluble in ether.

(a) To 2 c.c. aq. CuSO, [Febling's No. 1 solm is suitable] add dil. NH, OH until a clear blue solm is obtained, then dilute to 20 c.c. with water. To 5 c.c. of this solm add R.O. of O.S. and shake,

ppt. forma

Glycaryl tribenzoate, m.p. 76°.

(b) Priparation of haracectate, m.p. 119.
To 4 o. o. of socie subsytting in a t. t. add 2 drops cone. H.SO., Pour the mixture on to 1 g. of O.S. contained in a small beaker. After the violent reaction has ceased add 10 o.e. water and site. Filter off the solid, wash it well with water, crystallize from alcohol, dry, and determine the m.p.

DETERMINATION OF THE PERCENTAGE OF ACETYL RADICAL IN AN ALEXA ACETATE

Weigh out accurately into a 175-c.c. flask about 1 g. of the acctate.

Add 25 c.c. of approximately N. NaOH and boil gently for 15 min. Cool, add about 20 c.c. of distilled water and a few drops of phenolphthalein solm, then titrate the excess of alkali with N. HCl or H.SO.

Also titrate similarly 25 e.e. of the NaOH soln.

If v c.e. - the difference between the two titrations and w g. - the weight of acetate taken

then percentage of acetyl radical  $= \frac{43}{10} \times \frac{v \times factor of acid}{v}$ 

Alkonides, i.e. No or K derivatives of alcohols, dissolve in water,

forming strongly alkalms solutions owing to hydrolysis.

Mineral acids decompose the allroxides with liberation of the
alcohol, which may be distilled over, or separated, and tested by
the foregoing methods.

In any test where a mineral acid is employed, the alterride will give the same result as the corresponding alcohol.

#### PHENOLIC COMPOUNDS

Them are interested by the surfacture enter t spaces 25, 27). The phenotic character should be construct, when possible, by the surfacement of a characteristic derivative, such as an enter or an other,

Rots.

\$-Ketonie setere and \$-diketones behave like phonols.

Providers for the bloomfrature of O.S. :-

Under "OR liquid," or "OS, solid," in the appropriate table below, refer to the colour which has been obtained in the Fedtest with an appearus or alcoholic sein, and proceed as infinited. In certain cases the obour or lack of obour, and the solubility in

water of the substance, will need to be taken into account. If directed to pages 46-51, if "Monohythic phenols" or "Polyhydric phenols" or "Polyhydric phenols" or "Polyhydric phenols", apply any confirmatory tests, including a determination of mp., if OS, is a non-hyproscopic solid, given under the name of the phenol suspected. Also prepare and determine the mp. of one of the derivative there indicated, the first methods being usually the most suitable. The methods of preparation of the derivatives are given on caree 62-53.

O.S. Ilquid.
(4) Colour obtained by the addition of an FeCI, to an aqueous

Colone the of 0.5. Blue-violet " Carbolia" odour See (page 46) See "Quaiscol" Blue. rapidly Tarry odour (page 47) changing to red brown, solution becomes turbid Apply "NaOH test " Red-violet Pronounced edour

(page 45)
Wine-red
Pleasant fruity odour Proceed as follows:

To 4 in layer of neutral copper acetate in a t, add 5 cs. cold water, shake well, then filter. To the filtrate add 6 drops of 0.S. and shake. If a green ppt. is obtained see "#.Ketonic ceters" (page 113), or if a blue not is formed see "#.Ketonic ceters" (page 10).

(b) Colour obtained by the additic

soln. of O.S.

soln. of O.S.

PRELIMINARY PROCEDURE 45		
Red violet	Properties of 0 S Propounced ode	ur Apply "NaOH test."
Blue	Odour of cloves	(below) See "Eugenol " (page 47)
Green (transient	) Odour recembl that of cloves Odour recembl that of thym	ing See "iso-Fugenol" (page 47) ing See "Carvacrol"
offit test (for liquids giving a rect-violet colour with sq. FeCl.). To j.in. layer of O.S. in a.t. s. dad 2.c. a.g. Nabli and shake. (a) an immediate dense white ppt. is obtained, O.S. is prob- ably an ester of salteyin socid. See "Exters of carboxylio- acid" (togg 100). (b) an intense yellow soln. is produced, see "Phenolio abla- bydes" (page 64). : ctild. I Colour obtained by the addition of aq. FeCl, to an equeue- soln. of O.S.		
Colour	Properties of O.S.	
Violet blue, or blue	"Curbolia" odaus	See "p-Cresol" (page "p-Cresol" 47)
Violet blue, or blue	Odour of vanilla Sparingly soluble	Apply "Carbonyl test" (page 46)
Violet blue, or blue	Readily soluble in water	See ("Resor.   (page   50)
Violet	Sparingly soluble)	Apply "Carboxyl test" (page 46)
Blue-black	Odourices	See {"Tennic acid"} (page ("Gallio acid") 51)
Blue, repidly changing to red	Readily soluble in water	See "Pyrogaliol" (page 50)
Blue, instantly changing to red-brown	Readily soluble in water	See " Quinol " (page 51)
the, rapidly changing to red brown.	Tarry edour	See "Guaiscol" (page 47)
led-violet		Apply "Carbonyl Lest"

Apply "Carbonyl test" (page 46) See "Catechol" (page

49)

irren (changed Readily soluble in to red with a water drop of NaOH)

11) Tham received by the ablition of an Fell, to an akololic min of 0 s

0.5 Ral Thing Fracingly entable Phone whiles

Circum

Orem Cains and transient)

Apply "Carbenyl test" (below) See " a Naphthol " (page 49) Fee " \$ Nephthol " (page 42) Others of thymn Sen "Thymn!" (page 43)

Trata for additional groups in phenolic compounds:

Carborri un.

To J.in. layer of O.S. in a t.t. add an equal bulk of solid NaIICO, and 5 c.c. of cold water. Close the mouth of the t.t. and shake vigorously for I min., allowing any evolved gas to escape. Filter if any solid is present and carefully acidify the soln, or filtrate with cone, HCL

If (a) a ppt, is obtained, this indicates the presence of a carboxy? group in the molecule of O.S. Determine the m.p. of O.S.

and see Section 4 (page 79).

(b) there is no ppt, determine the m.p. of O.S. and refer to the list of m.p.s of monohydric phenols (page 47).

Carbonyl test. To 2 c.c. of dil. HCl add R.G. of 2: 4 dinitrophenylhydratine, . heat until the solid has dissolved, then cool. Add R.G. of O.S. and shake for a minute or so. If an orange or red ppt, is obtained see "Phenolic aldehydes" (page 64), or if there is no ppt. proceed as under "O.S. solid " (page 107)

MONOHYDRIC PHENOLS

Liquids. (See also solids of low m.n.)

"Carbolic" odour. Sinks in water.

(a) Apply Test (a) under "o-Cresol" (page 48) when a result similar to that given by phenol and o-crosol

will be obtained. (b) Apply Test (b) under "o-Cresol" (page 48), -blue violet colour (difference from phenol and

o-cresol which give a red colour). Tribromo derivative, m.p. 84°.

2:4-Dinitrophenyl ether, m.p. 714.

"Tohianasilohonate, m D. 50".

OH

B.p.

```
OCH, Tarry odour. Sinks in water.
205° Guaiacol.
          See "m.p. 28" Gusiscol " under "Solids."
                    OH
 237° Carracrol.
                          Odour resembling that of thyme.
                            Floats on water.
                  CH(CH.).
                  លអ
                    O-CH, Odour of cloves. Sinks in water,
 252° Eugenol.
                  ČH, CH:CH,
          Benzoate, m.p. 69°.
                    OH
                      OCH,
                               Odour resembling that of
                                 clores. Sinks in water.
                    ČII-CII-CH.
           Acetate, mp. 79°. Benzoate, mp. 103°.
Solids.
```

Bensate, m.p. 57°, p.Tolorosetybonate, m.p. 55°

OH Bp.

31° o-Creek OH 191°

\*\*Carlake " odom.

CH,

CH,

\*\*Thered CH, OH 182°

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M.p.

с, н, ют

(a) To R.G. of O.S. in a dry t.t. add a trace of solid

sodium nitrite, then add 5 drops cone. H.SO. Rotate the tube to mix the contents. A red colour indicates that O.S. is p.cresol. (Benzoate, mp. 71°. p.Toluenesulphonate, m.p. 69°. 2:4 Dinitrophenyl ether, m.p. 93°.1

A green or blue colour indicates that O.S. is phenol or o cresol. To the green or blue mixture add about 5 drops of water, then add aq. NaOH until the soln, is alkaline, -red colour on the addition of water and a blue or green colour on making alkaline (Liebermann's reaction).

- Apply Test (b). (b) In a dry t.t. place R.G. of phthalic anhydride and twice the bulk of O.S. Add 2 drops cone, H.SO. and gently heat until the mixture is red-brown in colour. Cool, add a few drops of water, then add aq. NaOH gradually with shaking until the mixture is alkaline. A red colour (due to the formation of phenolphthalein or its methyl derivative) indicates that O.S. is phenol or o-cresol. Distinguish by Test (c).
- (c) To } c.c. of molten O.S. add 2} c.c. conc. NH,OH and shake. Complete miscibility with the NH OH indicates that O.S. is phenol. (Benzoate, m.p. 68°. Tribromo derivative, m.p. 93°. p-Toluenosulphonate, m.p. 95°. 2:4-Dinitrophenyl ether, m.p. 69°.)

If O.S. does not dissolve in the NH OII this indicates that it is o-cresol. (2:4-Dinitrophe ether, m.p. 90°. Dibromo derivative, m p. ! p-Toluenesulphonyl derivative, mp. 53°. B zoate is liquid.)

CH. ČН(СП.)..

Dissolve R G. of O.S. in 2 c c, warm 50% acetic aci cool Add an equal shake, -violet red en p.Toluenesulphonate, m

Errer France Fra

Brakes States St

بجع

asi asi M.p. OH

Faint odour

aphthol. OH | (a) To 1-in, layer of O.S. in a t.t. add 2 c.c. aq. NaOH

and one drop of chloroform; warm. A blue colour is obtained with both a and finaphthols. Distinguish by Test (b). (b) Add R.G. of OS, to 10 cc. of a mixture of equal volumes of iodins soln, and aq. NaOH; shake. A riold colour, rapidly darkening, followed later

by a ppt. indicates that O.S. is α-naphthol. (Figure, m.p. 183°, p-Tolorersulphomato m.p. 89°,)

No change indicates that O.S. is β-naphthol. (Actite, m.p. 70°, Benzone, m.p. 100°, Pro-

(accrate, mp. 70°. Benzoate, mp. 107°. Piorate, mp. 156°. p-Toloenemiphomate, mp. 125°)

#### POLYHYDRIC PHENOLS

Solids.

General Properties.

(1) Readily soluble in water.

(2) Easily exidised, shown as follows :-

(a) Add 1 e.e. of the sq soin, of O.S. to Tollen's reagent (I e.e. sq AgNO<sub>2</sub>, I e.e. sq. NaOH; add dl. NH,OH drop by drop until a char, colouries soin, is just obtained)

A prey or brown ppt is given immediately by esteched, quinol, pyropalled, paller evid, and tannie evid; within 2 me, by rerend and phlorogludinel; and within 2 min. by reservant.

(I) To him layer of O.S. in a ti, all fice, an NiOH and shake. Owing to existing by the air property, tanget and, paths and, and grind pull dark red ferom solutions, other phenols pull training property or feroms additions. Nin. Oll.

on

Frank and a

C. 11. 101 M.D.

OH

To 2 e.c. of the sq. min. of OS. add an equal relume of had arreate,-immediate white ppt.

Diaretate, m.p. 63°. Dibenzoate, m.p. 84°. CH. Orcinol. Hydrated, m.p. 58

Resortinol.

(a) To 1 c.c. of the ag, soln, of O.S. add 1 c.c. aq. NaOH and one drop of chloroform. Heat the mixture and when a definite red or violet-red colour is obtained fill up the t.t. with water. A red soln, appearing rioles red on shaking, yielding no finorescence on diluting indicates that O.S. is resorcinol. Confirm the identity of O.S.

by applying Test (6). A riolet red soln, changing to deep red, yielding a green fluorescence on diluting indicates that O.S. is orcinol. (Yields no fluorescence in Test

(b)). Dibenzoate, m.p. 88°. (b) In a dry t.t. place R.G. of phthalic anhydride and twice this bulk of O.S. and then add 2 drops cone. H.SO4. Gently heat until the mixture is a red-brown colour. Cool, add a few drops of water, then add aq. NaOH gradually with shaking until the mixture is alkaline. Pour 1 c.c. of this alkaline soln, into a t.t. and fill up with water. A yellow-green fluorescence indicates that O.S. is resorginol. Dibenzoate, m.p. 117°.

OH OH 132° Pyrogallol. Orloweless.

- (a) Apply Tests (a) and (b) under "General pro-
- perties" (page 49). (b) To 2 c.c. of the ag. soln. of O.S. add 2 c.c. aq. lead acetate,-pale yellow emulsion changing to a
- heavy white ppt. in about 1 min. (c) To 2 c.c. of the aq. soln. of O.S. add a trace of powdered FeSO, and shake,-blue-violet rolour. Triacetate, m.p. 161°.



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THE PHENOIS IN THE FOREGOING LISTS (See sections on "Crystallisation" and "Drving of substances,"

pages 16-21.)

#### Acetates.

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In a dry t.t. place 1 c.c. or 1 g. of O.S., add a mixture of 21 c.c. sectic anhydride and one drop conc. H.SO, and shake for I min. Pour into 10 c.c. water contained in a small beaker and stir until the oil solidifies. (If the contents of the t.t. set solid, as may occur

during the acetylation of quinol or of pyrogallol, add water and stir). Filter, wash the solid well with cold water, crystallise from a mixture of 2 pts. alcohol and 1 pt. water (use acctone for pyrogallol triacetate), dry, and determine the m.p.

## Benzoates.

In a 100-c.c. flask dissolve 1 c.c. or 1 g. of O.S. in 5 c.c. acctone (see note 2), then add 21 c.c. benzoyl chloride. Add 50 c.c. aq. NaOH (the first 10 c.c. or so gradually with cooling and shaking, then the remainder all at once); cool. Cork the flask and shale vigorously for 10 min. (If time permits continue the shaking until the odour of the benzoyl chloride has practically disappeared.) Filter, wash the solid, first with dil. HCl, then with cold water, crystallise from alcohol (see note 1), dry, and determine the mp.

- (1) In the case of the benzoates of resoreinol and β-naphthol, if is is desired to crystallise the whole of the derivative, acctone will be found to be a more suitable solvent, since a large quantity of alcohol would be required.
  - The benzoates of quinol and phloroglucinol are only sparingly soluble in alcohol and in acctone; benzene is a suitable
- solvent for these derivatives. (2) By the use of acetone a cleaner product is obtained and the phenol, especially if solul, is brought into more intimate contact with the acid chloride.

# p. Toluenesulphonates.

In a 100-ca, flack place I ca. or I g. of O S., 2 g of p tolumns. sulphonyl chloride and 5 c.s. acetone. Heat on a water bath until all the solid has dissolved, then excl. Add 20 co. aq. NaOH, orth the fink and shake vigorously for 10 mm. Filter, wash the solid with sold water and return it to the 100 ce. flash. Attl 30 ce. as NaOH and some prevan per, best to healing and continue belling gently for 10 min. with free points shaking sound. (By this procedure the excess of sulphonyt chierds is seavertest buts the solution and an other is and solium p telementiphones, the arri supplement

ester being practically unaffected.) Add about an equal volume of water, cool and abake until the derivative solidifies. Filter and wash the solid well with cold water.

If, from the results of tests, O.S. is assumed to be

(a) phenol, gusiscol, thymol, α-naphthol, or β-naphthol, crystallise the derivative from alcohol.

(b) one of the cresols, dissolve the derivative in petroleum ether, filter into a small beaker and blow air (by means of bellows) over the surface of the liquid until a sufficient quantity of the derivative crystallies out.

Dry, and determine the m.p.

Bromo-derivatives of phenol, o-cresol, and m-cresol.

Dissolve 1 c.c. of melten O.S. in 5 c.c. sectons, and pour the sola. into a 100-c. of Reds. Add strong Br sola. 10 c.c. Br, 15 g. Rhr. 100 c.c. waterly until after chalding, the liquid in pale yellow. (About 2 c.c. will be required for c-resol, and about 18 c.c. for m-cresol, or phenol.) Add 60 c.c. water, cool and abake vigorously. Filter, was the solid well with end water, crystallis from a mixture of 2 pts. alcohol and 1 pt. water, dry, and determine the m p. Picrates of the maphthols.

Dissolve j-in. layer of 0.8. in a t.t. in 2 a.c. benzens. To the cold soln, add 2 a.c. of a saturated soln, of pierie acid in benzens and shake. Filter, carefully wash the sold with a few drops of benzens, dry rapidly by placing a thin layer on filter paper and waving over a small fishme, and determine the m.p.

2:4-Dinitrophenyl ethers.

In a 100-a.c. wide-monthed flask place 1 g, of O.S., 1 g, of C.S., 1 g, of C.S. and InterOnderstance, 10 c. a. c., an (Na-CO, 20 c. c., alcohol and some porous pot. Fit the flask with a reflux condenser, heat the contents to boiling and continue boiling for 10 min. Add 30-40 c.c. water, cool and shake. Filter off the solid, wash it well with odd water, crystallies from alcohol, dry, and determine the mp.

#### PHENOXIDES

Phonoxides (phenskes) dissolve in water and owing to hydrolysis yield strongly alkalino solutions. On passing CO<sub>4</sub> through the soln, or more rapidly by seldifying with dil. HCl, the phenol is liberated and may be filtered off, or extracted with ether (according to its solubility in water) and examined by the foregoing methods.

# ALDEHYDES AND KETONES

Included with the ablehydes are polymers and acctals which yield ablehydes on briling with mineral acids. Samples of polymers and acctals containing a free ablehyde may be succuntered.

# Liquide. O.S. completely miscible with twice its volume of water.

A. Akichydes and acctals detected by Schiff's reagent (Test 26, page 20).

Procedure for the identification of acetaldehyde, methylal, and formalin.

- (i) To R.O. of 2: 4-dinitrophenylhydrazine in a t.t. add 2 c.c. of dil. HCl and heat until the solid has dissolved. Cool the soln. and add j c.c. of O.S. With actaldhyde or formalin an immediate yellow ppt. of a 2:4-dinitrophenylhydrazone will be obtained, whereas with pure methylal a yellow ppt. (2:4-dinitrophenylhydrazone of formaldebyde) will be formed only after standing for a minute or two, or on warming the mixture.
  - for a minute or two, or on warming the minute.

    The presence of a carbonyl group in the molecule of O.S. or one of its products of hydrolysis is thus indicated.
  - (ii) Add one drop of O S. to Tollen's reagent (I c., of sq. AgNOs, I c.c. of aq. NaOH; add dil. NH<sub>2</sub>OH drop by drop with shaking until a clear, colouriess soln. is just obtained).
    An immediate gree or brown ppt. or a mirror of Ag
    - An immediate grey or brown ppt, or a mirror of Ag will be bitained with acetaldehyde or formalin but not with pure methylal.
  - (iii) Apply Test (a) under "Acetaldehyde." If the soln. remains clear and colourless, apply Test (a) under "Formalin" and distinguish between methylal and formalin by odour and b.p.

B.p. Oisagreeable, suffociting 21\*. Acctaldehyde. CH<sub>3</sub>-CHO. Disagreeable, suffociting odour (in dilute sq. soln, the odour resembles that of apples).

(a) To 2 c.c. of the prepared aq. soln. of O.S. add 2 c.c.

of 20% aq. KOH, heat to boiling and continue polling for 1 min .- soln. becomes yellow and a vellow ppt., changing to orange, with a disagreeable odour, is obtained,

55

16) To 2 c.c. of the prepared sq. solm. of O.S. side 2 c.c. of 1% aq. sodium nitroprusside, then add 5 drops of aq. NaOH-deep wine red colour. 2. 4-Dinitrophenylhydrazone, m.p. 168°. Semicarb-

szone, m.p. 162°. 42° Methylai. Ethereal odonr.

В.р.

Aprily Test (a) under "Formalin" (below) when a similar result will be obtained as the cone. H.SO.

first hydrolyses the methylal to formaldehyde and methyl alcohol. 97°-98° Formalin (Commercial 40% aq. soln. of formaldehyde,

H-CHO). Pungent odour. On evaporation leaves a white residue of paraformaldehyde, (CH.O)n. (a) To 2 c.c. of the prepared aq. soln. of O.S. add R.G. of resorcinol, then pour 2 c.c. of conc. H.SO.

(contained in another t.t.) carefully down the eide of the tube-red ring at the junction of the liquids; white ppt., which changes to violet red, forms in the aq. layer. (b) Oxidation to formic scid, II-COOH.

To 2 c.c. of the prepared aq. soln. of O.S. add 2 c.c. of H.O. (20 vol.) and a drop of phenolphthalein soln., then add so, NaOH, drop by drop with shaking, until a persistent red colour is just obtained. Stand the t.t. in boiling water for 5 min., cont, and again add an, NaOU, drop by drop with shaking, until a persistent red colour is just ob tained. Add dil. HCl until the red colour is jus removed, then add I e.e. of aq. HgCl. heat to

boiling and continue boiling for } min. -white ppt, of Hg.Cl. (due to the reduction of

the HgCl, by the formic seid produced). 2:4-Dinitrophenylhydrazone, m p. 166°. B. Ketones indicated by the red colour obtained with aq sodium pitroprasside and sq. NaOH (Test 26, page 27). All the priones in this section give a positive result in the

## following test :--

ladeform reaction.

To 2 e.c. of iodine soin, add one drop of O.S., then add ag

56 C. H. [0] NaOH, drop by drop with shaking, until the deep brown colour disappears,—pale yellow ppt. of iodoform, with characteristic odour. Procedure for the identification of O.S. Determine the b.p. of O.S. and refer to the following list of

П

b.p.s of ketones. If one of these b.p.s is identical with, or near to, that of O.S., confirm the presence of a carbonyl group in the molecule and the identity of O.S. by preparing and determining the m.p. of the derivatives there indicated. For methods of preparation of derivatives, see page 66. B.p. 56° Acetone, CH, CO-CH,

2:4-Dinitrophenylhydrazone, m.p. 126°. Semicarbazone, m.p. 187°. 80° Methyl ethyl ketone. CH, CO-CH, CH, 2:4-Dinitrophenylhydrazone, m.p. 111°. Semicarbazone, m.p. 148°. 164° Diacetone alcohol. (CH<sub>2</sub>),C(OH)-CH, CO-CH<sub>2</sub>. Aq. soln.

neutral; yields acctone on distillation with sq. Into a 50 c.c. distilling flask pour 1 c.c. of O.S. and 20 c.c. of approx. 2N. NaOH and add some porous pot. Cork the flask and attach it to a

water condenser. Distil, using a t.t. as the receiver, until 6-7 c.c. of distillate is obtained. From the distillate prepare acctone 2:4-dinitrophenylhydrazone, m.p. 126°. 165° Pyruvic acid. CH CO-COOH. Aq. soln. strongly acid. In the nitroprusside + NaOH test (page 27,) a violet-red colour is obtained, changed to violet with acetic acid; this latter colour quickly changes (a) To 2 c.c. of aq. KMnO, add one drop of O.S. to yellow.

4

-purple colour changes almost immediately to (b) To 2 c.c. of bromine water add one drop of OB. -immediate decolourisation. (c) In a t.t. place R.G. of β-naphthol, one slrop of O.S. and 2 e.e. of cone, H.SO.; shake,

-red colour becoming violet-blue on warming 2: 4.Dinitrophenylhydrazone, m p. 213°.

185" Acctonyl acctoms. CH, CO CH, CH, CO CH, 2: 4. Tanitrophenythydraums, m p. 118". Semioarb O.R. not misrible with senter

Proceed according to whether O.S. floats on water (below) or sinks in water (page 61).

C. O.S. floats on water. Add I or 2 drops of O.S. to 3 c.c. of Schiff's reagent and shake more or less continuously for 2 min. Do not heat the mixture. A deep red or deep violet colour indicates that O.S. is an aldehyde, or a polymer or acetal which has yielded an aldehyde by depolymerisation or hydrolysis. Some polymers and acetals, however, will not give a positive result, while with ketones the liquid will remain colourless or only a faint red colour will be produced. If a deep red or deep violet colour is obtained, confirm the aldehydia nature of O.S. by adding one drop of it to Tollen's reagent (1 c.c. aq. AgNO., 1 c.c. aq. NaOH: add dil. NH.OH. drop by drop. until a clear, colourless soln, is just obtained) and shaking for a period of time not exceeding 15 sec. A grey or brown ppt., or a mirror of Ag will be formed immediately or within 15 sec.

To identify O.S. determine its b.p. and refer to the following list of b.p.s of aldehydes, polymers of aldehydes, acetals, and ketones. If one of these b.p.s is identical with, or near to, that of O.S., confirm the identity of O.S. by filtering off, crystallising (see page 66), drying, and determining the m.p. of the 2:4-dinitro-

phenylhydrazone prepared in Test 2 (page 27).

For further confirmation of identity prepare and determine the m.p. of any other derivative indicated, or follow the given procedure. (For methods of preparation of derivatives, see page 66.)

B.p. 49° Propionaldehyde. CH CH CHO. Odour resembling that of acetaldehyde.

To 2 c.c. of the prepared ac. soln. of O.S. add 2 c.c. of 20% ag. KOH. Heat to boiling and continue boiling for 1 min ,-slight white ppt., dissolving to a clear pale yellow soln,; disagreeable odonr, (Difference from acetaldehyde which vields an orange ppt.)

2:4-Dinitrophenylhydrazone, m.p. 155°.

63° iso Butyraldehyde. (CH4), CH-CHO. 2:4-Dinitrophenylhydrazone, m.p. 182°.

64" Dimethylacetal. CH, CH(OCH,),

(a) Proceed as indicated under " Acetal " (b.p. 104") when similar results will be obtained owing to the hydrolysis of O.S. to acetaldehyde and methyl Andools.

(b) Proceed as indicated under "Acetal" but using

20 c.c. of dichromate mixture instead of the 20 c.c. B.D. of dil. H.SO4. To 2 c.c. of the distillate add a drop of phenolphthalein soln., then add aq. NaOH drop by drop with shaking until a permanent rel colour is obtained. Just remove the red colour by adding dil. HCl, add I c.c. of aq. HgCl, heat to boiling and continue boiling for I min, -white ppt. of Hg (Cl), due to the reduction of the HgCl by the formic seid produced by the exidation of the methyl alcohol liberated by hydrolysis of O.S.

C, H, [0]

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Yields the 2:4-dinitrophenylhydrazone of acetaklehyde, m.p. 168°. 74° n-Butyraldehyde, CH CH CH CH, CHO. 2:4-Dinitrophenylhydrazone, m.p. 122°.

80° Methyl ethyl ketone. CH, CO CH, CH, 2: 4-Dinitrophenylhydrazone, m.p. 111. Semicarbazone, m.p. 148°.

88° Ethylal, CH (OEt).

To 2 c.c. of water in a t.t. add one drop of OS, and R.O. of resorcinol, then pour 2 e s. of cone. II,50, (from another t.t.) carefully down the skle of the tube, -red ring at the junction of the liquids; white ppt , which changes to violet red, forms in

posults) and othyl alcohol.

the act soin. The cone, H 80 a first by indyses O.S to formaldehyde (detected by the above

a distantamenth of exame of formakle.

B.p. 2:4-Dinitrophenylhydrazone, m.p. 190°. Semicarbazone, m.p. 198°.

104 Acetal. CH, CH(CEt)<sub>2</sub>. Odour not unpleasant; does not affect the eyes.

- Into a 60-a.c distilling flash your 1 ac. of O.S. and 10 a. of Gil. II.504 and said some porous pot. Cork the flash and attach it to a water condenser. Distil, using a t.t. as the receiver, until 6-7 ac. of similizate is obtained. Text the distillate for acetaldehyde (produced together with ethyl alcohol by hydrolysis) as follows:—
  - (a) To 2 c.c. add 2 c.c. of 20% aq. KOH, heat to beiling and continue beiling for \( \frac{1}{2}\) min.,—yellow soln., followed by a yellow ppt., changing to orange: disagreeable odom.
  - (b) To 2 c.c. add 2 c.c. of 1% aq. sodium nitroprusside, then add 5 drops of aq. NaOH,—wine-red colour. Yields the 2:4-dinitrophenylhydrazone of acetalde

hyde, m.p. 168°. 124° Paraldehyde. (C.H.O).

Proceed as described above under "Acetal" when similar results will be obtained as the distillation with dil. H.SO, converts paraldehyds into acetaldehyds.

Yields the 2:4-dinitrophenylhydrazone of scetalde-

hyde, m.p. 163°.

124° Di kepropyl ketone. (CH<sub>3</sub>)<sub>2</sub>CH-CO-CH(CH<sub>3</sub>)<sub>2</sub>.

2:4-Dinitrophenylhydrazone, m.n. 93°. Semicarb-

azone, m.p. 157.

(a) Apply the lodoform reaction (page 55) when a positive result will be obtained.

- (b) To 1-in. layer in a t.t. of a soln. of Br in CC2, add 2 drops of CS —immediate decolorisation, due to the presence of a double bond in the molecule of CS.
- 2:4-Dinitrophenylhydrazone (red: crystallise from glacial acetic), m.p. 200°. Semicarbazone, m.p. 164°.

130° cyclo-Protanone CH<sub>e</sub>-CH<sub>e</sub>

2:4-Dinkrophenylkydramos (yellow), m.p. 145°, Semiourbanese, m.p. 203° decomp. B.p. 139°

Acetylacetone. CH. CO CH. CO CH. Aq. soln. gives a red-brown colour with sq. FeCl. Gives a blue ppt. with neutral copper acetate soln.

Ŋ

(a) Apply Test 2 b (page 27) when a wine-red colour

will be obtained. (b) Distil I c.c. of O.S. with 20 c.c. of approx. 2N. NaOH and collect 6-7 c.o. of distillate. From the distillate (consisting of aqueous acetone) prepare

acetone 2:4-dinitrophenylhydrazone, m.p. 126, by Method 1 (page 66). 151° Methyl n-amyl ketone. CH. CO [CH.] CH. 2:4-Dinitrophenylhydrazone, m.p. 70°. Semicarb-

azone, m.p. 124°. 154° n-Heptaldehyde (Oenanthol). CH. [CH.]. CHO. Us-

2: 4-Dinitrophenylhydrazone, m.p. 106°. Semicarbpleasant odour.

azone, m.p. 109°.

155° cyclo-Hexanone. CH CH CH CU

2:4-Dinitrophenylhydrazone, m.p. 160°. Semicarbazone, m.p. 166°.

Oxidation to adipic acid, m.p. 150°. Into a 100 c.c. flask pour 1 c.c. of O.S. and 30 c.c. of dichromate mixture, heat to boiling and continue boiling gently for 5 min. Cool, filter and carefully wash the solid free from green chromium salt with cold water (adipio acid is moderately soluble in cold water). Crystallise the solid from

water, dry, and determine the m.p. 172° Methyl a hexyl ketone. CH, CO [CH,], CH, 2: 4 Dinitrophenylhydrazone, m.p. 67°. Semicarb

azone, m.p. 123° 204° d Citronellal. C.H.r. CHO. Odour of lemon rind.

Semicarbazone, m.p. 81°, I Menthone. C. H.O. Odour of peppermint.

2: 4. Dmitrophenylhydrazono, m p. 146°. 229° Citral, C.H .. CHO. Odour of lemon.

The natural ablehyde is a mixture of two geometrical japenera.

2: 4 Dantropheny Dydrazog 200° d.Carvone, Calliato. Other of .

D. O.S. sinks in water.

Determine the bp. of O.S. and refer to the following list of h.p.s of addrydes and ketones. If one of these h.p.s is identical with, or near to, that of O.S. confirm the identity of O.S. by preparing and determining the m.p. of one or more of the derivatives there indicated. (For preparation of derivatives, see page 69).

B.p.

161° Furfaral. CH:CH

CH.CH Odour somewhat resembling bitter almonds. Colourless when pure; becomes brown with age.

CHO

(a) Dip the end of a roll of filter paper into a mixture of equal volumes of glacial sortic acid and aniline and remove any excess of the mixture by pressing the end of the roll between filter paper. Boil a mixture of one drop of O.S. and 2 c., of water in st.t. and hold the end of the roll in the vapours, —paper turned dopr red.

(b) Add 2 drops of O.S. to 1 e.e. of Fehling's soln. (equal volumes of No. 1 and No. 2) and boil for

min., - red ppt. of Cu.O.

Phenylbydrazone, m.p. 97\*. Semicarbazone, m.p. 202\*.

178° Benzaldehyde, C.H. CHO. Odour of bitter almonds.
(a) Preparation of beazons, C.H. CH(OH)-CO-C.H.
m.n. 131°.

Dissolve a piece of KCN, about the size of a pee, in 1 c. of water in a t.t. Add 3 c.c. of alcohol and 10 drops of O.S. Stand the t.t. for 5 min. in about 200 c.c. of water, which has been heated to behing and from which the fame has been removed. Cool, add 5 c.c. of water, shake and filter. Weath the solid with water, crystallise

it twice from alcohol, dry, and determine the m.p. (b) Preparation of discussionstone, (C.H.CH.CH), OO, m.p. 112\*. (Pale yellow instrous plates).

Into a bolim-tube pour I ex, of O.S., 5 drays of acrons, 6.z. of alrobel, and 2 r. r. day, No.No. Heat the mixture to bolim, continue boling for I min, then cool and slake vigorously. Add 20 cm of water; shake and filter. Wash the yellow abld with cold water, crystaline it twice from alsohol, dry, and determine the mp.

C, II, [0]

B.p. Phenylhydrazone, m.p. 156°. Semicarbazone, m.p. 214°.

193° Phenylacetaldehydo. C.H. CH. CHO.

2: 4-Dinitrophenylhydrazone, m.p. 121°. Semicarbazone, m.p. 153°.

202° Acetophenone. C.II. CO-CII.

Add a drop of O.S. to 2 c.c. of 1% 4q. soln. of sodium nitroprusside, then add 2 drops of aq. NaOH,
—wino-red colour, turned blue on acidifying with acctic acid.

Semicarbazone, m.p. 198°. Phenylhydrazone, m.p. 103° (turns brown and shrinks at 100°).

210° Propiophenone. C<sub>2</sub>H<sub>3</sub>·CO·CH<sub>3</sub>·CH<sub>4</sub>. Somicarbazone, m.p. 174°. 2:4-Dinitrophenylhydrazone, m.p. 191°.

225° p-Methylacetophenone. CH CO-CH.

Apply the test under "Acetophenone" (page 62) when a similar result will be obtained. Semicarbazone, m.p. 205°.

247° Cinnamaldehyde. C<sub>6</sub>H<sub>1</sub>·CH·CHO-CHO. Odour of cinnamen. Phenylhydrazone (yellow), m.p. 163°. Semicarbazone, m.p. 215°.

248° Anisaldehyde CHO

OCH.

Phenylhydrazone (white), m.p. 120°. Semicarbaione, m.p. 203°.

Oxidation to anisis acid, m.p. 181.

In a 250-ac. flask place the equivalent of j-linlayer in a t.t. of solik Khūto, and 60 a. of water. Heat to boiling, allow to go just off the boil and add 1 cc. of O.S. Gently abake round (it is not necessary to heat further until the colour of the Khūto, has disappeared; ccol. Filter, and scidify the filtrate with cone. Id. P. Biter of the solid, wash it with water, crystallise from alcohol, dry, and determine the m.p.

Calle

Solids.

Proceed as indicated below under E unless a yellow ppt, was obtained in Test 25 (page 30), in which case ascertain first if 0.8. is one of the commounds described under F (page 64).

Determine the m.p. of O.S. and refer to the following list of pa of aldebydes and ketones. If one of these m.p.a is identical, th, or near to, that of O.S. confirm the identity of O.S. by ring and determining the m.p. of one or more of the derivatives are indicated. (The methods of preparation of the derivatives not given under the name of the aldebyde or ketone, will be and on pages 65-66.)

d.p.
37° Piperonal. CH

(a) Preparation of arime, m.p. 110°.

Dissolve § g. of O.S. in 2 cc. of alcohol in a st. and da a soln. of § g. of hydroxylamine hydrochloride and 1 g. of crystallised column acetate in 2 cc. of water. Stand the st. for 10 min. in about 200 cc. of water which has been heated to biding and from which the fiame has been removed. Cod, fifter of the solid, wash it with cold water, crystallise from dilute alcohol, dry, and determine the m.p.

(b) Orndation to piperonylic acid, m.p. 228°.
Oxidise the equivalent of 4-in, layer of O.S. in a t.t. in the manner described for "Annialdehyde" (page 62). Czytallien the acid obtained from diute alcohol, dry, and determine the m.p. Pocnylhydrazone, m.p. 100°. Semicarbazone, m.p.

Bensophenone (Diphenyl Letone). C.H. OO-C.H.

Preparation of arise, mp. 1017.
In a 100-cc. wide-monthed flask place § g. of hydrorylamine bydrochloride, 1 g. of ands KOHI, and 5 cc. of vater. Add a sola. of § g. of O.S. in 5 cc. of alcohol and some person pot. Fit the flask with a reflar water condrawar, best the contents to boiling and continue boiling gently for far it, is better pikel of critice will be obtained if the boiling is continued for 1 kr.). Cool, add 20 cc. of water and slake. Filter from unchanged lettone and add to the filtrate d.l. HOI mittle better the pixel or critical in the pixel water and slake. Filter from unchanged lettone and add to the filtrate d.l. HOI mittle better and the given the pixel water and the pixel the pixel of the pixel or the pixel water and the pixel of the pixel or the pixel of the pixel of

Penyllydrame, mp. 137.

64 3( p.

Deoxybenzoin, Calla CH CO-Calla

2:4-Dinitrophenylhydrazone, m.p. 195°. Phenyl-p-tolyl ketone. C.H. CO-C.H. CH. 2:4-Dinitrophenylbydrazone, m.p. 199°.

134° Benzoin. C.H. CH(OH) CO-C.H.

(a) To 1-in, layer of O.S. in a t.t. add I c.c. of Fehling's soln. (equal volumes of No. 1 and No. 2) and I c.c. of water. Heat to boiling and continue boiling

for | min.,

-red ppt. of Cu O. (b) Oxidation to benzil, C.H. CO-CO-C.H., m.p. 95. To | g. of O.S. in a dry t.t. add 1.25 c.c. of conc. HNO, (a larger volume of HNO, may cause nitration) and stand the t.t. in gently boiling water for 10 min. Cool, add 10 c.c. of water and shake until the red oil solidifies. Filter off the solid, wash it with cold water, crystallise twice from alcohol, dry, and determine the m.p.

Acetate, m.p. 83°, see "Vanillin triacetate," page 85)

179° d.Camphor. C.H.O. 2: 4-Dinitrophenylhydrazone, m.p. 177°.

F. Solid polymers of aliphatic aldehydes. Paraformaldehyde. (CH,O)n. Pungent, fishy odour.

Apply the tests under "Formalin" (page 55) when similar results will be obtained.

Metaldehyde. (C.H.O)n.

Proceed as under "Acetal" (page 59) using the equivalent of 1-in. layer of O.S. in a t.t., when similar results will he obtained.

G. Phenolic aldehydes.

Liquids. B.p.

196° Salicylaldehyde. OH

CHO

To a solution of R.G. of 2: 4-dinitrophenylhydrazine of alcohol add 2 drops of cone. H,80, -orange ppt, indicating the presence of a carbonyl group in the molecule of O.S.

Phenylhydrazone (greenish-yellow), m.p. 142°; semicarbazone, m.p. 229° (see page 68).

Solids.

Odour of vanilla

Preparation of triacetate, m.p. 88. To 1-in. layer of O.S. in a dry t.t. add a mixture of

4 c.c. of scetic anhydride and 2 drops of cone, H.SO. Shake until all the solid has dissolved. then add 10 e.e. of water and cool. Allow to stand, shaking periodically, until the oil solidifies. Filter off the solid, wash it with cold water, crystallise from alcohol, dry, and determine the m.p. Semicarbazone, m p. 229° (see page 86).

CHO 115° p.Hydroxybenzaldehyde.

See "Vanillin trucctate" (page 65).

Preparation of derivatives. (See also pages 16-21.)

# Phenylbydrazonea.

# O.S. Liould.

To 2 e.e. of 50% sectic soid aid 5 drops of phenylbydratine, best just to braing, add 5 drops of O.S. and shake. Add 5 cc. of water, abake vicenesty and real. Fitter, etc., as under "O.S. add"

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To 1 in. layer of O.S. in a dry t.t. add 1 c.c. of glacial scetic acid and heat until solution is complete. Add 5 drops of plenylhydrazine and heat just to boiling, then cool.

If no solid separates, scrape the glass in contact with the liquid with a glass rod in order to assist precipitation. When solid has separated add 5 c.c. of water and stir. Filter off the solid phenylhydrazone, wash it with cold water and crystallise twice from alcohol (or aqueous alcohol). Dry a thin layer on filter paper over a small flame and determine the m.p. (Many phenylhydrazones decompose on prolonged heating in a steam oven, or on keeping.)

# 2:4-Dinitrophenylhydrazones.

METHOD 1. For aldehydes and ketones miscible with water. To 1-in, layer of 2:4-dinitrophenylhydrazine in a dry t.t. add 10 c.c. of dil. HCl and heat until all the solid has dissolved. Cool the soln., add i c.c. of O.S. and shake. Filter off the solid, wash

it with cold water, crystallise from alcohol, dry, and determine the m.p. METHOD 2. For aldehydes and ketones not miscible with water.

Follow the procedure under "O.S. not miscible with water" (page 27) if O.S. is a liquid or if O.S. is a solid proceed as indicated in Test 2 c (page 30). Filter off the ppt., wash it with

cold alcohol, crystallise from alcohol (or glacial acetic scid if sparingly soluble in alcohol), dry, and determine the m.p. Semicarbazones. In a dry t.t. place 1-in. layer each of semicarbazide hydrochloride

and powdered crystallised sodium acetate. Add 1 c.c. of water and heat until the solid has dissolved, then add i c.c. of O.S. and shake. If O.S. bas

(a) mixed with the reagent, cool, filter off the solid, very carefully wash it with cold water, crystallise from methyl alcohol, dry, and determine the m.p.

(b) not mixed with the reagent, add I c.c. of alcohol.

If the contents of the t.t. (i) have set to a solid mass, add 5 c.c. or water and stir.

Filter off the solid, wash it with water, crystallise from alcohol (or glacial acetic acid if sparingly soluble in alcohol), dry, and determine the m.p.

(ii) have not set to a solid mass, warm until a clear soln. is obtained, adding if necessary more alcohol. Cool, add 5 c.c. of water and stir. Filter off the solid, wash it with cold water, crystallise from alcohol (or dilute

alcohol), dry, and determine the m.n.

### CARBOXYLIC ACIDS, Erc.

IMPORTANT GENERAL NOTES.

- Acid anhydrides and certain exters (acid esters, also methyl and ethyl formates, methyl lactate, methyl and ethyl oxalates) in addition to carboxylic acids, yield strongly acid appears solutions.
- (2) For certain tests an aqueous soin, of a neutral sait of an acid is required. Prepare as follows:—

To be layer of the soid in a t. add just sufficient dil Nil, Oll to yield an allatine soin, and dilute to 10 cc. with distilled water. Four the soin, into a dish and holl until a piece of red litmus apper momentarly interested in the soil. Is just no longer turned blue, keeping the robume between 5 cc. and to cc. by the addition, when necessary, of durified water. Finally good and make up the volume to 10 cc. with distilled water.

(3) The determination of the equivalent weight of an acid or anhydride is a valuable aid to identification. (For methods of determination are page 81.)

Procedure for the alcalification of O.S. —
If O.S. is a liquid follow the procedure given below; if a solid follow the procedure under "O.S. solid" (page 62).

### O.S. liquid.

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- If O.S.
  (i) is miscible with water, then to a sola, of one drop of O.S.
  In 2 c.r. of water add 3 drops of an, FeG., If a strong pulser color is predomed, see "Lartic and" (spec 73]; if no such soleur is obtained, apply the tests under "Formin add, etc." (spec 83) under the other of O.S. is Like that of rands better in which case see "a.Briyne acid" (spec 83).
  - (ii) thats on water, or sicks in water, determine the b p. and refer to the appropriate Let of b p.s. of compounds (page 19)
  - (a) is not identifiable as one of the compression in the fallowing loss proceed as indicated under "Escore of carboryte ands" (page 100).

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O.S. completely miscible with twice its volume of water. Louis et Br. Pungent odour. 4502. inn\* Formic acid. If COOIS 60-03. Miscible with

CH. COOH 118\* Acctic acid. 7105 CII. CII. COOII water. 140\* Propionie sciel. (a) To 2 c.c. of the neutral aq. soln. add an equal volume of aq. FeCl. A wine-red colour (viewed through the depth of the liquid), yielding a reddish

brown ppt, on boiling, is given by all three scids. Apply Test (b).

(b) To 2 e.c. of the neutral aq. soln. add 1 e.c. aq HgCl, heat to boiling and continue boiling for min. If no ppt, is obtained apply Test (c) A white ppt. (Hg,Cl,) indicates that O.S. is formic acid (or a formate). Apply the following confirmatory tests :-

(i) To 1-in. layer of O.S. in a dry t.t. add 5 drops cone. H.SO. Gently warm the mixture by rotating the end of the tube over a small flame and turn the mouth of the tube periodically to the flame. CO is evolved and burns with a characteristic blue flame.

(ii) To 2 c.c. of the neutral aq. soln. add an equal volume of aq. AgNO, and boil, -liquid becomes brown, then a grey ppt, of

silver appears. (c) To 2 c.c. of the neutral aq. soln. add 2 drops aq. FeCl, and then add I c.c. of amyl alcohol. Shake vigorously and allow to stand until the mixture separates into two layers. The transference of the red-brown colour from the aqueous to the alcohol layer indicates that O.S. is propionic acid (or a propionate). No coloration of the amyl alcohol indicates that O.S. is acetic said (or an acctate). Apply the following test for acctic acid (or an acetate) :- To i in. layer of O.S. in a dry t.t. add 10 drops absolute alcohol and 10 drops cone. II,SO4. Heat gently, cool and pour into 5 c.c. water contained in a dish-pleasant fruity odour of ethyl acetate.

162° n-Butyric acid. CH, CH, CH, COOH. Equiv. wt. 88-06. Odour of rancid butter.

### O.S. floats on water.

B.p.
155\* ise-Butyric scid. (CH\_),CH-COOH. Equir. wt. 88 00.
176\* ise-Valeric scid. (CH\_),CH-COOH. Equir. wt. 109-1.
186\* ar-Valeric scid. (CH\_),CH-CH\_COOH. Equir. wt. 109-1.
205\* n-Cappric scid. CH\_1(CH\_1),COOH. Equir. wt. 110-1.
225\* s.-Heptoic scid. CH\_1(CH\_1),COOH. Equir. wt. 140-1.
227\* n-Cappric scid. CH\_1(CH\_1),COOH. Equir. wt. 140-1.

237° n-Laprylic acid. CH<sub>1</sub>(CH<sub>2</sub>), COOH. Equiv. wt. 144-1. 253° Pelargonic acid. CH<sub>2</sub>(CH<sub>2</sub>), COOH. Equiv. wt. 158-1. Capric acid. CH<sub>2</sub>(CH<sub>2</sub>), COOH. Equiv. wt. 172-1. Oleo acid. CH<sub>2</sub>(CH<sub>2</sub>), CH : CH [CH<sub>2</sub>], COOH Usually

possesses a tallow-like odour. Equiv. wt. 282-3.
To i-in layer in a t.t. of a soln. of Br in CCl, add
one drop of O.S.,

—deep brown colour immediately removed, owing to the presence of a double bond in the molecule of the acid.

### O.S. sinks in water.

B.p. CH, CO
138° Acetic anhydride. O Pungent, irritating odour.
CH, CO,

Equiv. wt. 51-02. Converted into acetic acid by boiling with water.

Preparation of anilide, m.p. 114°, and p-toluidide, m p. 148°:-

In a dry boiling-tube place § c. of freshy distrible aniluse (or § g. p-boiling) and add 1 c. o. of U.S. When the reaction has easeed add 20 c. boiling water and boil until the lower layer has completely dissolved. In the case of the p-to-boiling in with dissolved. In the case of the p-to-boiling in with dissolved. (in the case of the p-to-boiling in with complex plants). (bof, fit case of the solid with cold water, dry, and determine the m p, CH.CH.C. CH.C.

163° Propionic anhydride. >O Pungent odour.

Equiv. wt. 65-04. Converted into propionic acid by boiling with water. Anilide, mp. 105°. p-Tolvidide, mp. 124°. (For

preparation, see under "Acetic anhydride.")

186° Diethyl oxalate. See "Esters" (page 100).

O.S. solid.

If O.S. is readily soluble in water, proceed as indicated under A:
if sparingly soluble as under B (page 74).

n C. H. [0] 70 (a) To 2 c.c. of the prepared aq. soln. of O.S. add a drop of Α.

phenolphthalein soln., then add  $\frac{N}{1}$  NaOH until after shaking

a red colour is just obtained. Heat just to boiling. (If the red colour disappears this indicates that O.S. is an ester.) Add half the volume of glacial acetic acid, heat to boiling, then add 2-3 drops of aq. CaCl,

If (i) no ppt. is obtained apply Test (b).

(ii) an immediate white ppt. is obtained this indicates that O.S. is oxalic acid, or, if the red colour disappeared on boiling, dimethyl oxalate (see below). Both substances give a positive result in the following test:-

To 1-in. layer of O.S. in a dry t.t. add 5 drops cone. H.SO. Gently warm the mixture by rotating the end of the tube over a small flame, and turn the mouth of the tube periodically to the flame. CO is evolved and burns

with a characteristic blue flame.

Oxalic acid. COOH COOH

Equiv. wt. 45 01 (anhydrous); 63 02 (+ 2H,0).

Dimethyl oxalate, COO CII.

COO CH., m p. 54°.

The methyl radical may be detected by proceeding as Indicated under " Details of method of hydrolysis" (page 100). Use 2 g. of O S. and 10 s.c. of 20% aq. KOH. Hydrolysis will be complete in 5 min.

(b) To 2 cc. of the prepared aq soln, of OS. (and also to an equal volume of an aq soin, of tarterio soid of similar strength) add 3 drops aq. FeCla.

If with the aq. soln. of OS, there is obtained

(i) a strong yellow colour (like that obtained with the tartario acki) with no ppt, proceed as indicated under

(u) a yellow ppt, a practically colourhos soin, or a reduch brown soln, proceed as indicated under Hection 1.

Fection 1.

Trees of the same in of 0.8 att an equal volume of ex-Proliminary Test K 11-11, wal state 1 7 1 min

L. to ministered man " Making short are he and and polyphylone" If the realis select

to be exampled become my "I make crated and and an 12 - 12 - 120 The

In either case determine the m.p. of O.S. and refer to the list of m.p.s of scids and anhydrides. If one of these m.p.s is identical with, or near to, that of OS, confirm the identity of O.S. by applying any tests given.

Saturated acids and anhydrides.

M.D. CH, COOH Glutaric acid. CH

Equiv. wt. 66-03. CH, COOH

Apply the test under "Succinic soid" when a similar

result will be obtained. p-Nitrobenzyl ester, m.p. 69\*. CH,-CO

119° Succinio anhydride, { Four, wt. 50-02. CII.CO

Apply the test under "Succinic said" when a similar result will be obtained. HOOO.

133° Malonio acid. Equiv. wt. 52-02.

> (a) Heat 2-in, layer of O.S. in a dry t.t. until the solid has melted and effervescence occurs (due to the evolution of CO. - sharp odour of sectio acid.

> (b) To 1-in. layer of O.S. in a dry t.t. add 2 c.c. acetic anhydride and boil -reddish vellow liquid with yellowish green flaorescence. p-Nitrobensyl ester, m.p. 85°,

CH.CH.COOH 150° Adipie acid.

Equiv. wt. 7304. CH CH COOH

Apply the test under "Succipie sold " .- violet red colour after rendering the mixture alkaline with NaOH.

Amide, m.p. 220° (see page 80). p. Nitroberayl seter. m.p. 106\*.

CH. COOH 185° Succinio acid. ( Equir. wt. 5902. CIL COOH

> Place R.G. of O.S. in a dry tt. and add twice the bulk of reservinol and 2 drops ores. H.SO. Cently best until the mixture is a red brown rolour. Cool, add a few drops of water, then add aq NaOli until the mirture is alkaline. Pour

> > ٠.

C. II. [01 72

I c.c. of the alkaline soln. into a t.t. and fill up with water,-yellow-green fluorescence. p-Nitrobenzyl ester, m.p. 89° (see page 232). Anilide, m p. 226° (see page 80).

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Unsaturated acids and anhydrides. M.v.

CH-CO >0 Equiv. wt. 49-01. Maleic anhydride. 11 CH CO

Apply the test under "Maleic acid" when a similar result will be obtained.

Crotonic acid. CH, CH: CH-COOH. Equiv. wt. 86-05. To 2 c.c. of Br water add 2 c.c. of the aq. soin. of 72\* O.S. and shake.

-soln, becomes colourless in about 5 sec. CH-COOH

Equiv. wt. 58-02. (Aq. solu. 130° Maleic acid. 11

CH-COOH gives no ppt. with aq. FeCl.) Only decolourises Br water on heating. Pour 2 c.c. Br water in each of two t.t.s. To one add an amount of O.S. equivalent to 1-in. layer in a t.t. Stand both tubes in boiling water for 2 min.

The liquid in the t.t. containing O.S. becomes colourless, while in the other t.t. the brown colour of the Br persists. p-Nitrobenzyl ester, m.p. 89°. 133° Furoic (Pyromucic) acid. CH : CH

Equiv. wt. 112-0. ĊH : C

COOH

(Aq. soln. gives a yellow ppt, with aq. FeCl.) To 2 c.c. Br water add 2 c.c. of the aq. soln. of 0 S. -soln, immediately becomes colourless.

p-Nitrobenzyl ester, m.p. 133° (see page 232). Section 2

a Hydroxy acids. Preliminary tests. (1) To 2 e.e. of the aq soln, of the free acid or salt and I ac.

Deniges soln., heat to boiling and seld sq. KMnO., drop by drop. If, on the disappearance of the purple colour, a white ppt. is obtained aco " Citric acid." (page 73). If no ppt is

(2) Place R.G. each of acid or salt and \$ naphthol in a dry t t. James H.SO., Immeree the end of the tt. in boiling water for \( \frac{1}{2} \) min., shaking the t.t. in order to mix the contents; cool.

If the colour obtained is

(a) green, see "Tartaric acid" (page 74).

(b) yellow-brown, or deep red-brown, see "Lactic acid" and "Glycollic acid" (page 73).

(c) bright yellow with a green fluorescence, see "Malic acid" (page 74).

(d) rine red, see " Mandelic acid " (rege 74).

18° dl-Lactic acid. CH<sub>2</sub>·CH(OH)-COOH. Usually a syrupy
liquid.

(a) To a drop of the acid, or j-in, layer of the salt, in a dry t.t. add 2 c.c. cone. H<sub>2</sub>SO<sub>4</sub>. Warm carefully just until the soln. becomes pain yellow. Cool and add 2 drops of a 5% alcoholic soln, of guaiscol.

guaiacoi,

--intense red colour (Glycolle seid and its salts
give a violet-red colour) Confirm by Test (b).

(b) To 1-in. layer of soid or salt in a dry ££, add 5 drops cone. H<sub>2</sub>SQ<sub>2</sub>. Warm, revolving the tube, just until the mixture becomes light brown. Cool, dilute to 5 cc. with water and add solid NaOH until the mixture is strongly alkaline. Heat to boiling and continue boiling for 1 mix.

—disagreeable odour of aidehyde resin.
79° Glycollie acad. CH\_OH-COOH. Equiv. wt. 78-03. Tends to delignesce.

(a) Apply Test (a) under "Lactic acid," when a violetred colour will be obtained. Confirm by Test (b).

(b) To jin, layer of the seid or salt in a £t, add an equal bolk of edid KMnO<sub>t</sub> and 10 c.c. aq Na<sub>2</sub>O<sub>2</sub>. Heat and shake mult the purple colour has entirely desappeared, then filter. To 2 c.c. of the filtrate add 1 c.c. of figurial acetic seid, best to beding and add 2-3 drops aq. CaCl<sub>2</sub>. —white pyt. (Ca craisely.)

CH. COOR

100' Citric acid (+ 11,0). COH)-COOH Equiv. wt. 7043.

CH COOH

(a) See "Preliminary test 1" (Denicle test), page 72.

(b) To 5 cc. of the neutral sols add 1 cc. aq CaCl, best to boiling, and continue boiling fy 1.2 mm.

C. H. (01 74 -heavy crystalline ppt. of Ca citrate. (On M.p. applying Test (d) under "Tartaric scid" to a soln, of a neutral citrate of an alkali metal, a similar result will be obtained. NH, citrate, however, merely yields a purple-brown soln.) CH(OH)-COOH Equiv. wt. 67-02, Hygro-LMalic acid. scopic. CH. COOH (a) See "Preliminary test 2, result (c)," page 72. (b) To R.G. each of the acid or salt and resorcinel in a dry t.t. add 2 drops conc. H,SO, Warm over a small flame just until effervescence commences. Cool, add a few drops of water, then add aq. NaOH until the mixture is alkaline. Dilute to 100 c.c. with water, -hlue fluorescence. 118° dl-Mandelic scid. C.H. CH(OH)-COOH. Equiv. wt.

-blue fluorescence.

118° dl-Mandelic seid. C.H., CH(OH)-COOH. Equir. wt.
152-1.

To \(\frac{1}{2}\)-in. layer of seid or salt in a t.t. add \(\delta\) e.c. of a
mixture of equal volumes of aq. KMnO, and dl.

H.SO<sub>4</sub>; heat,
—hitter-almond odour of benzaldehyde.
CH(OH)-COOH
d-Tartario acid. | Equiv. wt. 75-02-

CH(OH)-COOH.

(a) See "Preliminary test 2, result (a)," page 72.

(b) To 2 c. of the aq. soln. of soid or sait sdd. R. G. of powdered FeSO., shake to dissolve and sdd one drop of H.O. (20 vol.). Add 2 c.c. sq. NaOH, deep violet colour (Fenton's test).

(e) To 3 o.o. of the neutral soin, add 1 c.e. aq. CaCls and shake vigorously.

—heavy, crystalline ppt. of Ca tartrate (dissolved—by adding 1 c.e. glacial acetic acid and boilings).

by adding 1 c.c. glacial acetic acid and boints;

(d) To 6 c.c. aq. AgNO, add 3-5 drops of the neutral
soln., then add dil. MH,OH drop by drop with
shaking until the ppt, is just or almost dissolved.
Stand the 8.t. in boiling water,
—eliver mirror forms.

3. Procedure for the identification of O.S...

Determine the nump. of the add (see Note 1 p. 75) and poly the Fol, test described on p. 75. Refer, in the appropriate section as indicated by the Fol, test, to the heart of the process of the property of the property of the process of the property of the

near to, that of O.S., apply any tests given under that soid. Also, if further evidence of identity appears desirable, prepare and determine the m.p. of one of the derivatives there indicated. (The methods of preparation of the derivatives are given on page 80.)

FeG, test.

To 2 c. of an aq. soln. of a neutral salt of the acid (for preparation see Note 2; page 67) sold an equal volume, of aq. FeG; and note the colour of the solu, or of the ppt. obtained.

Section in which Bernle Inference that O.S. us described tht buff ppt. a carboxylio derivative of benzene or toluene (see Note 2) 1 (page 75) ddish brown ppt. an aliphatic dibasic soid llow ppt. cimpamie scid or anisie acid low soln, benralic acid ip violet colour salicylic acid, or acetylsalicylic acid (see Note 3) 4 (page 79) Also see Note Ubrown colour m-hydroxybenzoic ecid or

p-hydroxybenzoic acid 19.

rm. If Q.S. sublimes, or has not maked when the thermometer register, 2302, proceed as indicated under Section 2 (page 77). Capborylie derrativites of benames or tolkene yield the hydro-fatbon when heated with soda-lime, while hydroxybennois acids rield phenol. Proceed as follows—Mix a quantity of O.S. which would about half fill the both of an ignition tube with hree times its bulle of dry adel-lime. Introduce the mixture ato the tube and heat. Note if an odeur of phenol, or of enzene or tolkene, hy produced. (A more satisfactory method federeting the evolved benzene or tolkene is given on page 80), violet colour is obtained with acetylasticpts exict in the PCA, stowing to hydrolysis to salicytic acid during the preparation (the new page 10).

ingly soluble acids and anhydrides.

C,H,CO

Benzoic ambydride.

C,H,CO

Equiv. wt. 1130.

(a) Hydrolysis to betazoic acid, m.p. 121°. To 1-in. layer of O.S. in a t.t. add 2 c.c. aq. NaOH and heat until a clear soln, is obtained. Cool, acidify with cone, HCl and filter. Wash the solid with oold water, dry, and determine the mp

U

C. H. [0] 74 (b) Preparation of benzanilish, m.p. 163°. 31.p.

To 1-in. layer of O.S. in a dry t.t. add 1 ec. aniline. Stand the t.t. in boiling water for 10 min., then remove; add 10 c.c. dil HCl and shake. Filter, wash the solid with sold water,

dry, and determine the m.p. Hydrocinnamic acid. C.H. CH. CH. COOH. Equiv. wt. 48\*

Apply the test under "Phenylacetic scid," when a similar result will be obtained.

Palmitic acid. CH [CH 1] COOH. Equiv. wt. 2583 Amide, m.p. 105°. 620 Stearie acid. CH. [CH.] .. COOH. Phenylacetic acid. C.H. CH, COOH. Powerful perfume 69° 76°

In a boiling tube place the equivalent of I in layer in a t.t. of O.S. Add an equal bulk of solid KMnO, and 5 c.c. dil. H.SO, heat to boiling. -bitter-almond odour of benzaldehyde.

Amide, m.p. 157°. Anilide, m.p. 117°. p-Nitrobenzyl ester, m.p. 65°. COOH

CH. Equiv. wt. 136-1. 102° o-Tolnic acid.

Amide, m.p. 141°. p-Nitrobenzyl ester, m.p. 91°. COOH

CH. Equiv. wt. 136-1.

Amide, m.p. 94°. p-Nitrobenzyl ester, m.p. 87°. Benzoic scid. C.H. COOH. Equiv. wt. 1220. Amide, m.p. 123°. Anilide, m.p. 163°. p-Nitrobenzyl ester, m.p. 89".

31° Phthaliq anhydride. ( You Equiv. wt. 7402.

(a) In a dry t.t. place R G. of O.S. and twice the bulk of phenol. Add 2 drops cone. II,50, and gently heat just until the mixture is a red brown colour. Cool, add a few drops of water, then add aq. M.p. NaOH gradually with shaking until the mixture is alkaline,—red colour of phenolphthalein, destroyed by acid.

(b) In a dry t.t. place R.G. of O.S. and twice the bulk of resortinol. Add 2 drops cone. H.S.O. and gently heat just until the mixture is a red-brown colour. Cool, add a few drops of water, then add aq. NAOH until the mixture is alkalme. Pour 1 cc. of the alkalme mixture is alkalme.

up with water,—yellow-green fluorescence. p.Nitrobenzyl ester, m.p. 155.

CH, CH, COOH

150° Adiple acid. | Equiv. wt. 73-04.

Apply Test (b) under "Phthalic anhydride," (page 76)
-violet-red colour after rendering the mixture
alkaline with NaOH.

Amide, m.p. 220°. p-Nitrobenzyl ester, m.p. 106°.

178° p-Toluic scid. Equiv. wt. 1361.

ČH,
Amide, m.p. 158°. p-Nitrobenzyl ester, m.p. 104°.
CH, COOH

185° Succinic scid. | Equiv wt. 59 02.

Apply Test (6) under "Phthalic anhydrade" (page 76) when a similar result will be obtained. p-Nitrobenzyl ester, m.p. 88" Anlide, m.p. 226".

195° Phthalic acid. COOH Equir. wt. 83-02.

Apply Tests (a) and (b) under "Phthalic anhydride" (page 76) when similar results will be obtained. 'p-Nitrobenzyl ester, m p. 165°.

Section 2.

Determine the equivalent weight of O.S. by Method I (1928 81) and refer to the list of equivalent weights of saids below. It one of these equivalent weights is identical with, or near to, that of O.S. apply any test spirm. Also prepare the derivative there indicated and determine its mp. The method of preparation of the derivative is given on page 89.

```
C. H. [0]
Equir. wt.
                          CH-COOH
                                        n-Nitrobenzyl ester.
  53 02 Fumarie acid.
                                                   m.p. 151°.
                          CH COOM
             Apply the test under "Maleic acid" (page 72) when
                a similar result will be obtained.
                           11000
                                    COOH
           To 1-in, layer of O.S. in a t.t. add 2 c.c. dil. NH OH
               and shake until O.S. has completely dissolved.
               Add I c.c. aq. BaCl, and shake vigorously. A
               heavy white ppt. indicates that O.S. is terephthalic
               acid, the Ba salt of isophthalic acid not being
               pptd, under these conditions.
           Methyl isophthalate, m.p. 64°. Methyl terephthalate,
              m.p. 140°.
     Cinnamic acid, C.H. CH : CH-COOH, Equiv. wt. 149-1.
          To 2 c.c. of the neutral soln, add an equal volume of
            aq. KMnO4 and shake, -purple colour immediately
             destroyed; brown ppt, and bitter-almond odour
             of benzaldehyde produced.
          Amide, m.p. 147°, Anilide, m.p. 153°, p.Nitro-
             benzyl ester, m p. 117°.
100° Benzilio acid. (C.H.), C(OH) COOH. Equiv. wt. 228-1.
          To R.G. of O.S. add a drop of cone. H.SO, -intense
             red colour.
          p.Nitrobenzyl ester, m.p. 99°.
                    COOH
                           Equiv. wt. 152-1.
                 m p. 162°, Anilkie, m p. 171°, p-Nitrobensyl
```

78

Eaction 4.

M.p.

135°. Acetylsalicylic acid (Aspirin).

СООН Ососи,

Equiv. wt. 180-1.

 (a) To 2 v.c. of cold water add R.G. of O.S. and shake, then add one drop of aq. FeCl<sub>2</sub>,—no violet colour (distinction from salecylic acid).
 (b) Apply Tests (b) and (c) under "Salicylic acid"

) Apply Tests (b) and (c) under "Salicylic acid when similar results will be obtained.

HOOD

158° Salicylic acid. OH Equiv. wt. 138-0.

(a) Apply Test (a) under "Acetylsalicylic acid" when a violet colour will be obtained.
 (b) To 1-in. layer of O.S. (acid or salt) in a t.t. add

I ca. cone. HNO<sub>2</sub>. Stand the it. in boiling water for 5 min. then remove and fill up with water. To 2 ca. of the yellow sola. (picric acid) and aq. NaOH until alkaline, then add 2 drops NH, sulphide and stand the tt. in boiling water for I min.—deep red colour due to an alkali salt of picramic acid.

(c) To 1-io. layer of O.S. (soid or sait) in a dry t.t. add 10 drops each of methyl alcohol and come. HSO, and heat gently. Cool and pour into 6 cc. water contained in a dish,—odour of "Oil of wintergreen" (methyl salicylate).

COOH

200° m-Hydroxybenzoic acid. OH Equiv. wt. 1380.

Apply Test (b) under "Salicylic scid," when a similar result will be obtained. Acetyl derivative, m.p. 131".

Acetyl derivative, m.p. 131°. COOH

213° p-Hydroxybenzoic acid. Equiv. wt. 1390,

Apply Test (b) under "Salicylo acid," when a similar result will be obtained. Acetyl derivative, m.p. 187.



standing for a few seconds.) Filter, weak the solid with cold water, crystallise from methyl sloohol in the case of the terephthalate, or from aqueous methyl sloohol in the case of the isophthalate. Dry the solid and determine the m.p.

Acetyl derivatives of m-hydroxybenzolc acid and p-hydroxybenzolc acid.

To jin layer of the said in a t. add 2 c.c. actic amfyddio and one drop one. H.SO<sub>4</sub>; heat just to boiling. Cool, pour into 5 c.c. water and heat until the lower layer has completely dissolved. (If the lower layer almost immediately solidines, as will be the case when phytmarylenoic acid is accelptatel, no further heating is necessary.) Tool and shake, filter, wash the solid with cold water, dry, and determine the m.p.

p-Nitrobenzyl esters. See page 232.

DETERMINATION OF THE EQUIVALENT WEIGHT OF AN ACID OR ANHYDRIDE

(A list of equivalent weights is given on page 86.)

Maked 1

Titration with  $\frac{N}{10}$  NsOH, with phenolphthalein as indicator.

(e) Standardise an approximately NaOH soln. (4 g. NaOH per htre) as follows:—

Weigh accurately a watch-glass or wrighing bottle, add about 0.2 g. of pure suceinic acid (equiv. wt. = 5902) and weigh again. Wash the acid with dutilited water into a beaker (230 c.c. or larger) or through a sufficiently large funnel into a 200-ce. coriolal flask.

Add 2 or 3 drops of phenolphthakin soln, and titrate with the NaOH soln, until a red colour, which persists for a minute, is obtained.

(0.2000 g. of succinic acid requires 23.9 c.e. of exactly  $\frac{N}{10}$  NaOH )

Factor of alkali - wt. of acid × 1,000

Other pure organic acids may be used, e.g. ozalic (hydrated, equir. wt. = 6362), tartane (equir. wt. = 7362) or citrio (hydrated, equir. wt. = 7063).

(2) Wrigh out 02-0-6 g of the and or and white whose equir, wt. is to be determined. If readily soluble in cold water wash it into the healer or flask with dealled water; if partiarly while in whi water was neutralized should (prepared by

п

Preparation of benzene or bluene from their carbarylic derivatives. To 1-in, layer of O.S. in a dry t.t. add 1-in, layer of dry solslime, pour into a morter and grind together, then return the mixture to the t.t. Fit the t.t. with a cork and bent delivery tube (with the long limb about 7 in.), place the t.t. across a tripod and arrange a second dry t.t. so that the end of the delivery tube reaches to the bottom of it. Heat the mixture fairly strongly but not so strongly as to soften and spoil the t.t. Finally heat the whole t.t. and the upper part of the delivery tube so as to drive any liquid present into the receiver. To the distillate add 3 drops each of conc. HNO, and conc. H.SO, warm slightly, dilute to 5 cc. with water and pour into a dish. A bitter-almond odour of nitrobenzene or the somewhat similar odour of a mono-nitrotoluene indicates that the distillate is benzene or toluene.

METHODS OF PREPARATION OF THE DERIVATIVES INDICATED UNDER THE ACIDS IN THE FOREGOING LISTS

(See sections on "Crystallisation" and "Drying of substances," pages 16-21.)

Amides, anilides, and methyl esters.

(These preparations should be carried out in a fume cupboard. Also, when adding a reagent to the crude acid chloride, the t.t. containing the former should be held at arm's length.)

First convert the acid into its chloride as follows - In a porcelain dish place & g, of the acid and 2 g, PCI, Grind the substances together by means of a pestle until the mixture becomes liquid. (If O.S. is assumed to be isophthalic acid or terephthalic acid use. twice the amount of PCI, and assist the liquefaction by warming.)

Preparation of the amide.

To the crude acid chloride add 10 c.c. conc. NH.OH. When the vigorous reaction has ceased, stir, cool and filter. Wash the solid with cold water, crystallize from water, dry, and determine the m.p.

Preparation of the anilide.

Dissolve the crude acid chloride in 5 c.c. acetone and pour the soln, into a 100-c.c. flask. Add 1 c.c. freshly distilled aniline and cool. Add 30 c.c. aq. NaOH, cork the flask and shake for 10 min. Filter, wash the solid with cold water, crystallise from alcohol. dry, and determine the m p.

Preparation of the methyl ester of isophthalic acid, or

To the crude acid chloride add 10 e.e. methyl alcohol and stir. then add 10 ec. water. (If OS, is tereof the salt with 50% H.SO, using a dropping tube. Place the crucible on a pipe-clay triangle supported on a retort ring, and adjust the height on a retort stand so that when a Bunsen is placed underneath, the bottom of the crucible will be about 4 in. above the top of the burner.

Heat with a very small flame and gradually raise the temperature of the crucible by bringing the ring pearer to the flame. (The very gradual heating is necessary in order to prevent excessive frothing and the creeping of the mixture too far up the sides of the crucible.) When the contents are dry, place the crucible in an inclined position on the triangle, and heat by placing a small flame under the lower rim. Gradually move the flame along until it is directly under the bottom of the crucible. Finally heat to redness in a large flame until any carbon present has burnt away and the residue is white. (The removal of carbon is facilitated by using a large Meker burner and moving the crucible about.)

Allow to cool, moisten with 50% H.50, and drive off the excess aced in the same manner as before, finally heat to redness, cool in a desicestor and weigh.

Repeat heating, for periods of 10 min , until the weight is constant The percentage of No. K. Sr. and Pb. as well as Ba and Ca in salts

may be similarly estimated. The student should, for practice, carry out duplicate estimations of the recentage of metal in several commercially pure salts without removing any water of ervatallustion recent

Below are shown duplicate results obtained with commercially pure salts, together with the percentage of metal calculated from the formula :

3

C

i - 2	•	('air-intel	Frank Info.
Mrium	formate, (CHO <sub>4</sub> ) <sub>2</sub> Ca	% Ch = 20 62	30 23 0 26 2
•	bertate, (C,11,O,1,Ca,511,O	1200	13 23 0 13 %
-	terines, C.II.O.C.4II.O	35 40	13-43 0 21%
٠	Olitary (Ca)! "Util Carellio	51-07	2192 0195
-	parameter (Cl31011CF2H10	1:40	1247 010
	arrian, fCallanalan	2 14 - 40 ts	50 3C
in lum	brosumin Cally Oaks	% No - 13 97	1575 0 120



41 Possible diffusing from the thereshout factor by get more than mat, should be obtained

Paswet Pri til party e of he sinnamate riskled owns g of he is paint of Ag is contained in 0.779 g of Ag mit.

Espite wt of soil - 2551 - 1079 + 1 - 1492. .. calculated from formula - 149.1.

(2) DAZIO g. of Ag. benzests gave 0-2000 g. of Ag.

Equir. wt. of benzole scid = 122 2. (Theory 1220) Market 3

If the addition of cone, aq BaCl, or CaCl, to a cone, neutral Analysis of a Ba or Ca sait. soln, of the NH, sait of the acid yields a ppt., the Ba or Ca sait obtained (after filtering off, washing, and drying), may be used for the determination of the equir, wt. of the acid.

The disadvantage of the method is that the salt may contain water of crystallisation, and it is therefore necessary before use to heat it in an air oven to about 130° until constant in weight.

The method employed is to heat a weighed quantity of the Ba or Ca salt with H.SO, and to weigh the residue of BaSO, or OaSO. From the weight of sulphate obtained the weight of Ba or Ca in the original salt is calculated.

Thus  $\frac{137.4}{233.4} \times \text{wt. of BaSO}_4 = \text{wt. of Ba in the original salt.}$ 

$$\frac{40.08}{136.1}$$
 × wt. of CaSO<sub>4</sub> = ", ", Ca " "

Next the weight of salt equivalent to 68-68 g. of Be (or 20-04 g. of Ca) is calculated. From this is subtracted 63-68 g., for the gram equiv. wt. of Ba (or 20-04 g. for the gram equiv. wt. of Ca), and 1 g. is added for the H which the Ba (or Ca) has replaced; the result is the gram equiv. wt. of the acid.

For the heating it is convenient to employ a retort stand fitted Procedure :with a ring upon which rests a pipe-clay triangle. (An alternative method is to use an Argand burner in the manner described in method 2 (b)). Heat a crucible (preferably a deep silica one) to redness on a triangle, cool in a desiccator and weigh.

Place a layer of the Ba or Ca salt (1 in. to 1 in. deep) in the bottom of the crucible and weigh again. Just moisten the whole KOH and boil until all the solid has dissolved., (An effervescence will be observed, due to the evolution of oxygen.) Acidify the soln, with conc. HCl, cool and filter. Wash the solid with cold water, dry, and determine the mr.

### SALTS OF CARBOXYLIC ACIDS

stification of the acid radical.

only Na or K is present see "Procedure I"; if any other I is present see "Procedure II" (page 89).

where I:-

0.8.

- s) does not yield an acid aq. soln., to j.in. layer of O.S. in a t.t. add 5 c.c. distilled water and shake. If O.S. does not dissolve, heat until solution is complete, then cool.
  - If (i) a clear soln, is obtained, dilute to 10 e.e. with distilled water and proceed as under A.
    - (ii) a lather was produced during the dissolving of O.S. and a white gelatinous mass was obtained on cooling the soln, a sait of palmitic, stearic, or oleic acid is indicated. Proceed as under B (case S9).
  - ) yields an acid ag, soin, to j-in, layer of 0.8. in a 11, and efficient dif, NH, QH to yield an alkaline soin, and diste to 10 ee, with dutilied water. Four the soin, into a dist to 10 ee, with dutilied water. Four the soin, into a dist and boil until a piece of red titmes paper momentarily immersed in the soin is just no leager turned bloc, keeping the volume between 3 cc. and 10 cc. by the addition, when becomenty of dutilied water. Cool, make up the work of the control of the dutilied water. Only include the case of proceed as under A.
    - o 2 cn. of the neutral soln and if the volume of glacial vite axid. If a ppt, is obtained (undesting the liberation a sparingly soluble axid) apply Tert (b); if no ppt, is odned, heat to boiling and axid 2-3 drops ag. CAG, An immediate while ppt, inclusive as creatist; apply the

d under (ii), page 70.
If no ppr. is obtained apply Test (b).

2 c.c. of the neutral solt. (and also to an equal volume of ter as a blank test) ald 3 drops aq. Fe'll, observe the ct and then ald an equal volume of aq. Fe'll,

	Equivalent weights	of acids a	nd anhydrides
•	(Calculated—C = 12	00, H = 1	009, 0 = 1000
46 02. 49 01. 50 02. 51 02. 52 02. 58 02. 59 02. 60 03. 63 02. 66 03. 67 02. 70 03. 73 04. 74 02. 74 02.	Formie acid. Maleie anhydride. Succinio anhydride. Acetio anhydride. Maleio scid. Maleio scid. Maleio scid. Succinio scid. Succinio scid. Oralio scid. Oralio scid. Oralio scid. Oralio scid. Malio scid. Citrio scid. Malio scid. Citrio scid. Thibalie anhydride.	86 05. 89-05. 102-1. 112-0. 112-0. 122-0. 125-1. 136-1. 136-1. 150-1. 150-1. 125-1. 125-1. 225-1.	Butyrio acids. Lactio acids. Valerio acids.
83-02.	Phthalio scids.	284 3.	Diegra -

# LACTONES AND PEROXIDES

CII: CII M.D. 67° Coumarin.

Fragrant ordour. Sparingly soluble in water. Dissolves in hot aq NaOff yielding a yellow solu. which on anklifying with HCl gives a white pps. of coumarin acid.

Preparation of II fCl, compound, m p. 165. To 1 in layer of US in a 11 at 1 2 c c. alcohol, warm entil the solid has dissilved, then sool Aid a will win of \$ in. layer in a t t. of Hg(7, in 8 a.s. of 25", aboted; shake Filter off the ppt of fine nearlies which expectation wash with mile water,

dry, and determine the mp 127 Bennyl persada CH, COOOCUCH, Faint sweet colone familiatio in water Rapholes corresponding acid by method 2 (page 83), or method 3 (page 84) and refer to the list of equivalent weights of acids (page 86). If one of these equivalent weights is identical with, or near to, that of the acid of which O.S. is the sait, apply any tests given under that acid (pages 71–72).

B. Procedure for the identification of an alkali salt of palmitic, stearic, or oleic acid:—

Dissolve about 1 g, of 0.8, in 20 cc. bot water and acidify the soln, with cone, ICL. If the liberated said is a solid, heat until it has nelted.\* Four the hot mixture into a small separating funnel (containing about 10 cc. water to prevent exacting); cool. Extensively, with either (see page 21), dry the ethereal soln, with anhydrous CaCL, sidtill 60 the either and cool the residue.

If the residue is

(a) a liquid, see "Oleic scid" (page 69).

(6) a solid, see "Palmitic seid," and "Stearic seid " (page 76).

Procedure II:-

Pour 20 c.c. distilled water into a dish, heat to boiling and add the equivalent of \(\frac{1}{2}\). In a st. of anhytrous Na<sub>2</sub>O<sub>3</sub> and an equal bulk of O.S. Continue boiling gently with stirring for min. Test the soln, with red litmes paper; if not alkaline do sold Na<sub>2</sub>O<sub>3</sub> gradually with stirring until an alkaline reaction is 'obtained. Pilter.\*

Tour the clear filtrate (see note below) into a dish. If the volume, is considerably more than 6 a.e. buil down to about this volume. Brunove the fame and add did. HENO, with stirring until a piece of bibe lature spaper momentarily immersed in the son, is piece turned definitely red. Add did. NHO, OHI with stirring until any world which has been prick is completely dendered and the soin, is just alkaline. Buil until a piece of red litture spare momentarily immersed in the soin, is just to longer turned bibe, keeping the volume between 5 c.c. and 10 c.c. by the addition, when necessary, of datilide when the second of the soin is just and the soin in the soin in the soin is the soin to some the soin between 5 c.c. and 10 c.c. by the addition, when necessary, of datilide when the soin is just the soin to some the soin to some the soin the soin is the soin to some the soin that so the soin the

Ood, make up the volume to 10 e.c. with distribut water and apply the test under A (page 87). For Test (c) proceed as indicated above under "Procedure II" as far as the asterisk, but using about 3 times the amount of Na<sub>2</sub>O<sub>3</sub>, and of O.S. (see note). Ood the fittest, pour into a small backer, alt one. IICl with sturing with the legal is acid, then proceed as indicated in Test (c) after the aciditation with IICl.

NOTE.

If the fibrate is not perfectly clear, re filter until this condition is attained. If, however, the fibrate consents of a white emulsion,

Wath sound volume aq. FeCl. Wine-red colour (viewed through depth of liquid). No ppt. Reddishbrown ppt. on boiling. Deep yellow colour. No ppt.

Yellow ppt.

Reddish-yellow colour. Formate, acetate, or No ppt.

C. H. [0]

With 3 deops

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Informer and procedure

propionate. Apply

Test (b) under corre

sponding seids (rec

Proceed as indicated

under "a.hydroxy

Proceed as under (c).

Cinnamate or anisate

scids," (page 71). Salt of a phenolic acid

681.

Strong yellow colour. . Salt of a hydroxy scid. No ppt.

Violet or red colour. Deep violet colour.

Proceed as under (el. Salt of a simple salur. ated aromatio seel. Light buff ppt. Usually similar, le Prococel as under (c). Salt of an aliphate dense ppt. acid. Proceed as are Reddish-brown ppt. der (0). If results, other than those described above, are obtained, proceed as under (c).

(c) Prepare a soin of O.S. by dissolving about 5 times the amount of O.S. previously taken in the minimum amount of water-Add I ce cone. HCI, if no immediate ppt, cool and shake If (1) a ppt, is obtained, filter, wash the solul with cold vigorously.

water (see note), dry and proceed as indicated under B (page 74) If a pict was only obtained after centing and shak Nors ing, as would timisly be the even with a salt of secretain or arlipin acut, the washing must be carried

out very carefully (2) an estime of reacul butter, or other unphasens what ; or an oil to obtained, extract with other (see page 21) If my cal is present first esturate the order with earlydrain Caff. Dry the othereal with with anhydraus

Cally and distil off the other Intermine the be of the resulting west arm lack of the (pages an, 411) on mather of the position bearing make (1) and (2) to attacions increasing a true expensations weight of the

### CARBOHYDRATES

 Carbohydrates readily soluble in water, detected by Molisch's test. (Test 3, page 30.)

### NOTES.

(1) The aq soln, referred to in the tests is that which has been employed for Molisch's test, i.e. 1-in layer of O.S in a t.t. dissolved in 10 c.c. water.

(2) The m ps are of little value for identification purposes since carbohydrates seldom melt sharply owing to the fact that fusion is nearly always preceded by alight decomposition. Their specific rotations are the most uncertaint physical constants.

(For method of determination and values a suitable text-book should be consulted.)

Procedure for the identification of O.S .:-

To 2 cc. Fehing's soln. (equal volumes of No. 1 and No. 2)

add t cc. of the aq. soln. of O.S. and stand the t.t. in boiling water
for I mia.

If (a) the whole mixture becomes red, due to suspended Cu<sub>2</sub>O.

- proceed as under A.
- (b) the mixture remains blue, and is either clear, or only a very slight yellowish turbidity is present, proceed as under B (page 92).

### A. Apply the following test:-

To 2 et. of Parice's soln add an equal volume of the aq soln, of OS, and stand the tt. in briskly boiling water for 2 min. If a red ppt. of On 0 is observed in the blue soln, or on the sides of the tabe just above the hquid, apply the tests (preva lessor) for glacore, fructore, and glatcore; if no red ppt is obtained,

apply the tests for lactore and maltore (page 92).

Test for glaron, fracture, and galacton.

(b) To See of the say who, of OS. skil to drop of glarid screins and 3 drops of phenythylarance, skile. Lowely cert the 4t. and stand it is bolling water for 10 mm, persolately skiling. If a bully yriller yet, is produced, spriy Test (1); if no such per, no detaund, apply Test (2).

(The prilies ppt is an ensurer, the same con being formed from look glucose and frontese. The ensurer of galartise does not separate under the above conditions.)



formed by hydrolysis), indicates that O.S. is Sucrose (Cane sugar)  $C_{11}H_{11}O_{11}$ , m.p.  $160^\circ$ , or Inulia  $(C_1H_{11}O_4)n$  (decomposes on heating). Distinguish by Tests (b) and (c)

(b) To 1-in layer of O.S. in a t.t. add 2 c.c. aq NaOH, heat to boiling and continue boiling for 1 mm

A deep yellow soln, indicates that O.S. is Inulin. Sucrose

yields an almost colourkess soln.

(c) In a 100-c.c flask place the equivalent of a 1-in layer in a t.t.
of O.S. and add 50 c.c. dil. HCl (1 c.c conc HCl diluted to

100 c c. with water).
Heat to boiling, continue boiling gently for 10 mm, then cool.
Add a drop or two of phenolphthalein coln. and just sufficient

aq. NaOH to give a permanent red colour Add dil. scette acid, drep by drop, until the soln, is just colourless.

The soin obtained from either sucrose or inulin will reduce Fehling's soin, and also Barfoed's soin. (See under A, page 91).
The soin, obtained from sucrose, but not that from inulin, still give the test for glucose (b) (page 92) if double the amount

of lead acetate is employed.

Starch (C<sub>\*</sub>H<sub>10</sub>O<sub>\*</sub>)n.
 This carbohydrate will have been indicated by the blue colour obtained on the addition of iodine to its aqueous solution (Test (b) under "No metal present," page 29).

Confirm by the following tests—
(a) Apply Bloisch's test (Test 3, page 30). Use a soln obtained by boiling R.G. of O.S. with 2 c'c water and cooling.

A positive result will be obtained.

(5) Heat 50 c c. water, contained in a beaker, to boiling. Shake 1-in. layer of O.S. in a t.t. with 5 c.c. water, pour the muxture into the beaker, boil for 1 min., then cool. Apply the following tests:—

(1) Add 5 c.c. of the aq. soln. of O S. to 50 c.c. water to which one drop of iodine soln. has been added,

-deep blue colour.

Heat 5 c.c. of the blue soln, in a t.t.,—blue colour disappears and reappears on cooling.

(2) Add 2 cc. of the aq. soin. of O.S. to 2 cc. Fehling's soin. (equal volumes of No. 1 and No. 2), heat to boiling and continue boiling for i min.—no reduction, i.e. no red ppt. of Cu.O.

(c) Shake 1-in. layer of O.S. in a t.t. with 5 c.c. cold water.

Pour the mixture into 50 c c. oi boiling dil. HCl (5 c.c. conc. liCl diluted to 100 c.c. with water), boil gently for 10 min, then cool (a clear soln, will be obtained with potato starch whereas rice starch yields a slightly opalescent soln.). Add a

C. H. [0] (3) To 2 on of the art onto of O.S. will the equivalent of Im.

layer in a 11 of whit level arctate, heat to boiling and add See the NHOH Heat the mixture again to bolley and continue beiling for about 1 min. A salmon-pask of grown penk enhant inclinates that O.S. is Glidcone (Destrose,

Grape on gar) CH, OH [CH(OH)], CHO, m.p. 146 (ab) diora). arr per it mol 11.0) If a buff colour is obtained proceed with Test (c). (c) To 2 cc of the aq soln of O S add an equal volume of cone. HCl and R O of reservined Stand the t.t. in boiling fater for 2 min. A deep wine-red colour, usually followed by a ppt

which dissolves in alcohol yielding a deep wine-red solnindicates that OS is Fructose (Levulose, Fruit sugar) CII,OH [CII(OH)], CO-CH,OH, m p. 95°-105°. See also page 94. (d) Place 1 g. of O.S. in a dry t.t. Add 11 c.c. HNO. (con-HNO, + 1 its volume of water). Stand the t.t. in boiling water until effervescence commences and red fumes are evolved, then transfer the tube to water which has been heated to 70°. Allow to stand for 15 min., keeping the water at 65°-70°. A white, sandy ppt. (mucic acid) indicates that O.S. is Galactose, CH<sub>1</sub>OH [CH(OH)]<sub>1</sub> CHO, m.p. 168° (anhydrons) 119° (I mol. H.O).

(a) To 2 c.c. of the aq. soln. of OS. add the equivalent of a lin. layer in a t.t. of solid lead acetato, heat to boiling and add twice the volume of dil. NH OH (with lactose a dense white ppt, is obtained; with maltose only a slight ppt, is formed). Heat to boiling and continue boiling gently for I min.

A salmon-pink colour indicates that O.S. is Lactose (Maltos gives only an orange-yellow colour). Confirm by Test (b). (b) Proceed exactly as described in Test (d), page 92. Finally remove the t.t. from the hot water, cool and scrape the inside of the tube in contact with the liquid with a glass

rod for 3 min. A white, sandy ppt. (mucic acid) indicates that O.S. is Lactore (Milk sugar) C1H2,O11, m.p. 203° decomp. (The amount of ppt, will increase considerably on standing.) Maltose C, HarO11 (Malt sugar, m.p. 100° decomp.) does not

B. Apply the following tests for sucrose and inulin :-

(a) To 2 c.o. of the aq. soln, of O S, add an equal volume of cone. 1 D C of resorcinol. Stand the t.t. in boiling water

### COLOURED SOLID COMPOUNDS

Procedure :--

Determine the mp. of O.S., then refer, in the appropriate subsection, according to whether O.S. is yellow, carage, red, or green, to the last of mp s of compounds. If one of these mp as a stemic with, or near to, that of O.S., apply any tests given for that compound, and prepare the derivative undested. The method of preparation of phenylhydranones is given on page 63.

G.S. yellow.

At P.

41° Renzalacetone (Benzylidenescetone f
ketone) C.H.-CH:-CH-OO-CH.

fityryl methyl

(a) To f-in layer of a soln of Br in CCl, in a dry tt. add B.O of O.S and shake, —deep brown colour disappears almost instantly.

owing to the presence of a double bond in the molecule of O.S.

(b) Add R G of O.S to 1 e c of cold cone H<sub>c</sub>SO<sub>c</sub> and shale,
 —O.S dissolves yadding an crange-red soln.
 Phenythedranous, m.n. 150.

84° Placernone (Diphenylene ketone) Phenylhydrasone, m.p. 151°.



AN BOALL CH, COCKII,

In a dry pareclain data standing on a water both place a point of KOH, then place on the top of it 2 little of O.S. Carrilly point 1 s.s. of almost alreded dress the sale of the data, where is not within

Francista to digrees, remove any unfaced od EOH, all for even H.50, and prefly such the dak so that the and runs over the whole of the position. 91

sloop or two of phenolphthabin solo, and fast sufficient of NaOll to give a permanent red colour. Add dil. scetic scid, drop by drop, until the soln, is just colourless.

C, H. [0]

Report both the tests under (b) with this solu.

(1) No blue colour will be obtained with the lodine sola-

(2) Fehling's soln. will be reduced, i.e. the whole mixture will become red, due to suspended Ca,O. The above results are due to the fact that by boiling with the mineral seid the starch has been completely hydrolysed to

# glucor.

GLUCOSIDES

The only glucoside likely to be encountered is Salicin, C, HuO,

m.p. 2014. This substance will have been indicated in Test 3 (page 30) by the intense red colour given with cold conc. HrSO.

Confirmatory test. Boil the equivalent of 1-in. layer in a t.t. of O.S. with 25 c.c. dil. H.SO. (11 c.c. conc. H.SO. diluted to 100 c.c. with water) in a 100 c.c. flask.

In about 10 min. a white ppt, will appear (saliretan, a condensation product of salicyl alcohol). Pour I c.c. of the mixture into

t.t., add aq. NaOH until the mixture is alkaline, then add an equal volume of Fehling's soln. (equal volumes of No. 1 and No. 2) and boil for 1 min., -whole mixture becomes red, due to suspended Cu.O. (The reduction is caused by glucose formed by hydrolysis. An aq. soln. of salicin will not reduce Fehling's soln.)

To the mixture in the flask add the equivalent of 1 in. layer in a t.t. of solid K, Cr, O, and 10 c.c. dil. H, SO. Connect the fine to a water condenser and distil until 4-5 c.c. is present in the receiver. The distillate will possess the characteristic odour of salicylaldehyde. Apply the following tests for this aldehyde:-(a) To half the distillate add one drop of sq. FeCl, -riolet-red

colour. (b) To the other half of the distillate add one drop of ag. NaOH,

-yellow colour.

Further distinction between glucose and fructose: In a t.t. place 2. R.G. of O.S. and R.G. of sodium selenite (or SeO.J. Add 2 c.c. of dil. HCl and stand the t.t. in bolling water for 2 min. A finely-divided red ppt, of selenium is obtained with fructoso, whereas with glucose the soln, remains clear and colourless.

M.p.

285° Anthraquinone.

Insoluble in NaHSO, soln.

In a t.t. place 2 R.G. of O.S. and an equal bulk of zinc dust. Add 5 c.e. aq. NaOH, heat to beiling and continue boiling for a min.,

-deep red colour.

Filter while hot into a tt. and shake,
—red colour rapidly disappears, owing to exidation
by the air, and a floculent, pale yellow ppt. of
anthraquinons separates. The phenomena describer may be repeated by adding sine dust,

boiling, etc.

The red colour is due to the Na salt of exanthranol.

O.S. orange.

202° Phenanthraquinone



(a) To jun, layer of O.S. in a 21 add 3 er of satupated NallSO<sub>4</sub> soln, and warm the mixture, —O.S. dissolves, and is repyrd on the addition of any NiOll.
(b) In a 21, place 2 R.G. of O.S. and an equal htd.

e) on an insert access of the arm an equal total of size dost. Add Ser any North, best to bridge and continue bridge for § mm. Filter the red brown ada, which so obtained.

manist salta dag ad

1) and an analysis

zilio acid, (C.H.),C(OH) COOH. Diphenylhydrazone, m.p. 225°.

115° p-Benzoquinone.

Peculiar pungent odour. (a) To 2 c.c. KI soln, add one drop dil. H.SO, then

add R.G. of O.S., and shake, -deep brown colour, due to liberated iodine. (b) In a t.t. place the equivalent of 1-in, layer of

powdered FeSO, and an equal bulk of OS. Add 2 c.c. dil. H.SO, and shake. To the mixture add 5 c.c. water, and heat until a clear yellow soln. is obtained, then stand the t.t. in cold water. -quinhydrone separates in fine, long green needles with a metallic lustro.

(c) Preparation of phenominone (red), m p. 71°. Dissolve in separate t.t.s in 3 c.c. hot ligroin 1-in. layer of O.S. and 1-in, layer of phenol. Mis the two solutions and cool. Filter off the red needles which separate from the yellow soln, recrystallise from ligroin, dry, and determine the

m D.

Fungent odour, similar

to that of p-henzoquinone. Sparingly soluble in water, but dessolves in aq NaOH yielding a refbrown soln.

Call, CH(OH) CO Calls. Colourless when 130°+134° Benzoin pure, but yellow samples will often be encountered Ben " Benzoin " (page 62).

CH Colourkon when pure, but

> technical qualitum are yellow Box " Anthracene " (page 125).

Insoluble in NaHSO, soin.

In a t.t. place 2 R.G. of O.S. and an equal bulk of zine dust. Add 5 c.c. aq. NaOH, heat to boiling and continue boiling for 1 min.

-deep red colour. Filter while hot into a t.t. and shake,

-red colour rapidly disappears, owing to oxidation

by the air, and a flocculent, pale yellow ppt, of anthraquinone separates. The phenomena described may be repeated by adding zine dust, boiling, etc.

The red colour is due to the Na salt of exanthranel.

prenge.

Phenanthraquinone.



- (a) To 1-m. layer of OS, in a t.t. add 3 c.c. of saturated NaHSO, soln, and warm the mixture. -O.S. dissolves, and is repptd. on the addition of
- aq NaOH. (b) In a t.t. place 2 R.G. of O S, and an equal bulk of zine dust, Add 5 c.c sq. NaOH, heat to boiling and continue boiling for & min. Filter the red brown soln, which is obtained.

-blue-green residue on the filter paper.

đ.

Phenoquinone. C.H.O.2C.H.OH.

Place R.G. of O.S. in a porcelain dish. add a drop of dd. NH.OH and stir. -blue-green colour.

. 98 O.S. green. M.p. (1) 110. Readily soluble 171° Quinhydrone. C.H.O., C.H. in cold alcohol, yielding a yellow soin. Add R.G. of O.S. to Tollen's resgent (1 c.c. aq. AgNO. 1 c.c. aq. NaOH; add dil. NH,OH, drop by drop, until a clear colourless soln, is just obtained),

-immediate grey-brown ppt.

C, H, [0]

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# ESTERS, ETHERS, AND HYDROCARBONS

If O.S. is

(a) a liquid, completely miscible with twice its volume of water, ascertain if it is identical with "Dioxane" (page 114); if it is not proceed as indicated under "Esters of carboxylic acids" (page 190).

(The following esters are completely miscible with twice their volume of water—methyl lactate, ethyl lactate, ethyl tartrate.)

 (b) a solid, or a liquid, not completely muscible with twice its volume of water, proceed as follows:—

Weigh accurately a clean dry flask of 150 c.e. capacity. having a short wide neck and flat bottom. Add 1-1-5 g. of OS, and weigh again. Add 25 cc of approximately N. alcoholic KOH and 3 c.c. water. Fit the flask with a cork and reflux condenser, place on a water bath, and allow the contents to boil gently for | hr Meanwhile titrate 25 c.c. of the alcoholic KOH with N. HCl using phenolphthalein as the indicator. (N. H.SO, gives a ppt. of K.SO, with N. alcoholic KOH). When the contents of the flask have been heated for the required time, pour 10 c.c. water through the condenser into the flask. Also loosen the clamp holding the condenser and detach the latter from the flack, raising it so that the underside of the cork is about I in above the mouth of the flask. Wash, by means of a wash bottle, that portion of the inner tube which projects through the cork, so that the wash water falls into the flack. Cool the contents of the flack, aid phenolphthalein soin, and intrate the alkali present with N. HCL

Work out the value of z in the following expression

Wt. of O.S. × 1000
Difference between the two intrations × factor of acid

If the value of x is less than 500 proceed as indicated under "Esters of earliexyle acids" (page 100) etherwise see "Ethers and hydrorathons" (page 114).

Nore

The above procedure is constitutly that described on page 112 for the determination of the equivalent weight of an enter. The

14



boiling for a further period of about 1 of the time taken for the ester to disappear.

Notes.

(1) With esters of alcohols which are sparingly soluble in water there will always remain a layer of liquid on the surface of the ag. KOH. The majority of such esters are completely hydro-

lysed in 1 hr.

(2) Hydrolysis is accelerated by periodically loosening the condenser clamp and giving the flask a rotary movement.

Procedure after hydrolysis:-

Allow the contents of the flask to cool somewhat, then pour 10 c.c. water through the condenser. Disconnect the flask and attach it to a sloping condenser. If no liquid is floating on the surface of the aq. KOH proceed as under A (below), otherwise as under B (page 102).

A. Distil, using a t.t. sa the receiver, until 10 c.c. of distillate is obtained. (Ignore any slight milkiness which may be present in the distillate, due to a trace of unchanged insoluble ester passing over). Pour the alkaline residue in the flask into a beaker and keep it for the later treatment described on page 103. Rinse out the flask with water.

If the distillate possesses a pungent odour, resembling oil of mustard, apply the tests under "Allyl alcohol" (page 34), otherwise rour half the distillate into the empty flask which was used for the hydrolysis and add 20 c.c. dichromate mixture. (Keen the other half of the distillate in case of accident during the pext reocedure.)

If the dichromate is reduced (in which case the mixture will usually become green and hot within a min.) proceed as indi-

cated below under "Identification of the alcohol." If the dichromate is not reduced proceed as indicated under

"No alcohol has been detected " (once 104).

Identification of the alcohol.

To the contents of the flask add 2-3 pieces of porous pot, connect the flask to the aloping condenser and use a Lt. (marked to indicate the space occupied by 7 c.c.) as the receiver. Distil until 7 e.c. is present in the receiver, noting if, when 2-3 e.c. is present, there is a laver of oil on the surface of the distillate. Proceed as under A (page 32). When the alcohol has been identified, preceed to identify the said constituent of the ester in the manner described under "Treatment of the allaline residue to the beater " (rage 103).

100 C. H. [0]

n.

NOTE. In describing the procedure under (a) and (b), it is assumed for the sake of clearness, that the ester has been completely

hydrolysed and therefore the layer of liquid floating on the ad KOII is an alcohol. The possibility of incomplete hydrolysis and the procedure to be adopted in such a case, is dealt with under (c).

Distil, using a t.t. as the receiver, until about 3 c.c. of distillate is obtained.

If the distillate

(a) consists of two well-defined layers, continue distilling until the passage of oily drops through the condenser ceases. (The alcohol will form an upper layer, i.m. or more deep.) Pour the alkaline residue in the flask into a beaker and keep it

for future treatment. Insert a 10-c.c. pipette into the distillate so that its point rests lightly on the bottom of the t.t. Draw up the whole contents of the t.t. into the pipette. Wait for a few seconds

for all the small globules of alcohol to rise to the surface and unite, then allow the lower layer to run back into the t.t. Run the alcohol into a dry t.t., add about 1 of its bulk

of freshly ignited K<sub>1</sub>CO<sub>1</sub>, cork the tube and allow to stand for 1 hr. or more. Meanwhile proceed to identify the acid constituent of the ester in the manner described under "Treatment of the

alkaline residue in the beaker" (page 103). When the alcohol is dry, determine its b.p. and refer to

to the list of alcohols under "O.S. floats on water " (page 36). (b) consists of a white emulsion, together with oily drope, which sink in the water, transfer the distillate, together with the cooled contents of the distillation flask, to a small separating

funnel. Add 10 c.c. ether (see "Extraction with other, page 21), shake and allow to stand until two well-defined layers are formed. Run off the lower layer into a beaker and keep this alkaline residue for future treatment. Pour the ethereal soln, through the neck of the funnel into a dry t.t. and add about } of the bulk of freshly ignited K.CO. Cork the tube and allow to stand for | hr. or more. Meanwhile proceed to identify the acid constituent of the ester in the manner described under "Treatment of the alkaline

residue in the beaker" (page 103

the estimated with her sto

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pour it into a dry flask, (that used for the hydrolysis, after triasing out and drying, will be conveniently connect the fact to the water condenser and partly immerse it in hot water contained in a beaker. When all the ether has distilled to the residue apply Test (8) under "Benzyl alcohol" (page

2) is clear and no upper layer is vasible, collect about 15 c.o. of distillate. If there is found to be little or no upper layer in the t.t., or if the liquid (alluded to as "the alcohol") obtained by the procedure under (or of 0) cannot be identified as one of the alcohols under "O S. floats on water "(page 35) or as benzyl alcohol, momplete hydrolysis is inducated.

Repeat the hydrolysis as before but using 5 g, of solid KOH and 5 c, water instead of the 25 c n. of 20% an KOH. Boil rapidly, with frequent shaking, for 2 br. (Towards the end of the hydrolysis the contents of the flask may become semi-solid, owing to the separation of the K salt of the acid. Allow to cool somewhat, add 30 c.c. water through the condenser, then proceed as under B fores 103.

eatment of the alkaline rendue in the beaker.

Sool the liquid if it is not already cold. (If it is found that is contents of the beaker have set on a solid mans, due to the uaration on cooling of a sparingly soluble K salt, e.g. K manusto, reclaisolve the solid by carefully heating and add it sufficient water to prevent any solid separating again on oline.)

Then carry out the procedure under the appropriate heading, her immediately below, or on page 104.

s slcohol has been detected.

Addily 1 a.e. of the cooled alkaline soln, with a mixture of equal lumes of cone. INO, and water (corresponding to roughly 30% NO<sub>3</sub>), cool and shake. (This concentration of acid is used in der that organic acids, such as philadic acid, which are moderety soluble in water, may be pidd. No KNO, is deposited, era after cooling and shaking, when 20% aq. KOH is just acidified sha 30% KNO<sub>3</sub>.

If (a) a ppt, separates, aciddy the whole of the alkaline or raidue with 30% HNO, cool and shake. Filter off the pptd. organic acid, wash it with cold water and dry. Examine the soid in the manner described under B (page 74).

(b) no ppt. separates, proceed as under (1) if the acidified liquid possesses a rancid or unpleasant odour, otherwise as under (2). r. ft. for

\$11 A mility the whole of the alkaline Books with 30% HVD, and sald from! Call, until an ell separates. Ketract the nil in the manner described he the alcohol unifor 137 pages 102, but using freed CaCl, as the drying

agents. Intermine the by of the oil and refer to the list of b pa of liquid souls (page 68).

(2) Actify 2 o a. of the alkaline soln, with glacial acetic soul, heat to brilling and sail 2 or 3 drops of ar. Call; a white ppt, indicates an exalate.

If no white ppt, is obtained just acidly the whole of the alkaline soln, with 50% II,50, add cone. NH OH until just alkaline, cool and filter off any K.SO, which may have separated. Boil until the

soln is neutral, cool a portion and proceed as indicated under A. Test (5) page 87. If a solid salt is required for confirmatory tests evaporate some of the neutral soln, to dryness in a basin, heating over a gauze until spirting commences, then completing the removal of water by heating on a sandbath with rapid stirring. In tests, however, which involve warming with cone. H,50, eg. the guaiscol test for a lactate and the fluorescein test for a succinate, the original ester will give the same result

as the solid salt. If only negative results are obtained, add 10 c.c. of conc. NII,OH to 2 c.c. of the original ester and allow to stand for 1 hr. with periodic shaking. If a crystalline ppt. is formed, filter it off, wash it with cold water, dry, and determine the m.p., then refer to the list of m.p.s of acid amides (page 180) in order to identify the acid constituent of the ester.

No alcohol has been detected. The alkaline residue will contain a phenoxide, or a polyhydric

alcohol.

Procedure:-

Pour the alkaline residue into a 100-c.c. cylinder and make up to 50 c.c. with water. Pass a fairly rapid current of CO, (which has been washed with water) through the liquid until I c.e., after the addition of 10 c.e. aq. BaCl, gives no colour on adding phenolphthalein soln. (This indicates that no free KOH is present and therefore any original phenoxide, which would be partially hydrolysed by water to the free phenol and KOII, is completely decomposed.)

e co. for 15 min, is sufficient.

اغ بر احد If (a) a considerable amount of solid separates, filter it off (keep the alkalno filtrate), wash it with cold water to remove the K sait of the acid and K<sub>1</sub>CO<sub>2</sub> and dry. Apply Test 1 (page 29) and see "Phenolic compounds" (rare 44).

(page 44).

Extract the alkalune filtrate with ether, exactly as described under (b) in order to remove any phenol left in soln., which would interfere with the tests for acids, especially the FeCl, test. Identify the acid constituent of the cetter in the alkaline filtrate in the meanure described under "An alkaline filtrate with each of the second section of the meanure described under "An alkaline filtrate with each of the second section of the section of the

(page 103).

(b) little or no solid separates, or an emulsion is formed, transfer the liquid to a separating funnel, add 10 c.c. ether and shake. (See " Extraction with other," page 21). Allow to stand until two well-defined layers are formed, then run off the lower aq layer into a beaker. Pour the ethereal extract into another vessel, return the alkaline hould to the separating funnel, add 10 c.c. ether and proceed as before. Repeat the process a third time, leaving the ether soln, this time in the funnel and retain the alkaline liquid in the beaker. Pour the other two ether extracts into the funnel and allow to stand for a time, then run off any lower aqueous layer which separates. Dry the ether soln, by allowing it to stand in contact with freshly ignited K.CO, for as long a period as cottvenient, then pour the liquid through the neck of the funnel into a dry flask (that used for the hydrolysis is convenient) and attach the fissk to the water condenser. Distil off the other by partially immersing the flask in hot water contained in a beaker.

If a residue is obtained apply Test 1 (page 29), then see

"Phenolic compounds" (page 44).

Treatment of the alkaline residue in the beaker.

If (a) a phenol has been detected, proceed as indicated under
"An alcohol has been detected" (page 103), in order to

identify the acid constituent of the enter.

(b) no phenol has been detected, just acidity the whole of the
alkaline residue with 50%, H.50,, add cone, NH,OH until
just alkaline, cool and filter off any H.50, which may
have separated. Disappears to dryness in a basin and
cool. Six the solid or entail-acid residue with 10-15 co.

volume of ether; filter. Wash the residue on the filter
with 5 c. c. of the alcohol-ther mixture, allowing the

C, H, [0] 106 filtrate to run into the previous one. To the residus on

the filter apply Test (b) under A (page 87), in order to identify the acid constituent of the ester. Transfer the filtrate to a dry flask (that used for the bydrolysis is convenient), attach the flask to the water condenser and partially immerse the former in boiling water contained in a beaker. When the alcohol and ether have distilled off, examine the syrupy residue (which will usually contain solid matter) for the presence of ethylene glycol or glycerol (see pages 41, 42).

ſΙ

15

15

30

8

01

60

30

In the following lists, the time required for the complete hydrolysis of some of the commoner liquid esters (using 20% sq. KOH,

or equal weights of KOH and water) is given.

216° Ethyl succinate

234° Ethyl salicylate

2820 Methyl pathalate 294° Ethyl mirate .

-nes Fibri philalate

218° Methyl phenylacetate

258° Glycerol triacetate (Triacetin) .

224° Methyl salsoylate .

209 Ethyl phenylacotate

271° Ethyl cinnamate . 250° Ethyl tartrete

Of educa aciento or and							
Hydrolysis with 20%	aq.	кон					
Esters, of which the al	anh ati	c con	stituen	t is n	niecibl	e wi	V Some
Esters, of which the un	CONLOS						Minutes
B.p.							5
32° Methyl formate			•		•	- 1	15
54° Ethyl formate		•		•	•		15
57° Methyl acetate		•		•	•		15
77° Ethyl acetate			•				10
79° Methyl propiona	te.		•		•		30
91° 40 Propyl acetat	ο.			•	•		20
98° Ethyl propionate			•		•		20
101° n-Propyl acetate				•	•		25
2020 Mathyl n.butyrat	ο.			•	•		30
190° Ethyl n-butyrate				•	•		10
145° Methyl lactate			•	•	•		10
esse Ethyl Inciate .	-		•	•	•		10
121° Methyl maionate		•		•	•		5
rose Ethel ovalate.		. •			•		10
1970 Fibriege glycol d	aceta	ute	•	•	•		10
195° Mothy! succinate	•		•	•			15
198° Ethyl malonate		•		•			20
198º Methyl benzoate	•	•	•	•			45
213° Ethyl benzoate		•		:			20

٠

# \* ESTERS OF CARBOXYLIC ACIDS 10 Esters of which the alcoholic constituent is not miscible with water

· - -

B.p.							
98° iso-Butyl formate	•		-				I hr.
107° n.Butyl formate							**
116° iso-Butyl acetate							
124° iso-Amyl formate							
126° n-Butyl scetate							
139° iso Amyl acetate			- 1				
188° n-Butyl lactate	-						
203° Benzyl formate						-	
216° Benryl acetate	•	:	:	:	:		
228° Benzyl propionate	•	:	:	:	:		39
268° n-Butyl saheylate							**
277° iso-Amyi salicylate			•			-	**
211 Mo-Amys saucylate	٠.	•	•	•	•		*1
320° Benzyl salicylate	٠		•	•	•		**
1	Phe	nolic i	alers				
B.p.							Monutes
196° Phenyl acetate					-		5
278° Resorcinel disceta-	te						10
Hydrolysis with equ	ıal	weig	hts	of K	OH a	nd 1	rater
B.p.							
241° iso-Butyl benzoate	•						1-2 hr.
249° n-Butyl benzoate							
262° iso Amyl bensoate							-

#### O.S. solid

Procedure:-

318° Benzyl phenylacetate .
323° Benzyl benzoste .
338° n-Butyl phthelate .
349° iso-Amyl phthelate .

Determine the mp. of O.S. and apply the following test:— To j.in. layer of O.S. in s t.t. add an amount of solid KOH roughly equal in bulk to a pea and 2 c.c. alcohol. Heat to builing and continue builing for j min.

If a dark brown or green colour develops proceed as indicated under C (page 109); otherwise as under A if m.p. 80° or below, or under B (page 109) if m.p. above 80°.

A. Pils 100-c. flack, having a short wide neck and flat looton, with a cost and a reflat a condenser. Benove the flack from the condenser and place in it 5 g. solid KOH, 5 c.e. water, 2 g. of 0.5, and sone provus pot. Connect the flack again to the condenser, beat the contents to boiling and continue boiling for j hr. undees the ester completely disappears in a shorter period of time. (Some methyl esters and also some phenolin esters will be completely hybridy-seld in 5 min. or lead.)

c. n. (9)

t-a

If (a) no vil is present on the surface of the eq. ROH. allow the contents of the fisk to cool sometim well 30 e a water thereigh the enderer, then proces servily as described under A (page 101).

Alms pefor to Bat A (page 110), in order to secrate thy taking into account the mp. of OS) the sleebel or phenol and acki likely to be present and proced accordingly If, on account of the mp, and the proved absence of a phenoxide in the alkalme residue. is is eneported that OS, is either ethylene girel dibenzoate or glyceryl tribenzoate, the procedure described under (b) page 105, should be modified as

Acklify the alkaline reskine with a mixture of equal follows :volumes of conc. HNO, and water; cool. Filter off the pptd, acid and apply tests for benzoic acid. To the filtrate add cone, NH, OH until just all shre. Evaporate, etc., as described under (b) page 105.

(b) an oil is present on the surface of the sq. KOH, allow the contents of the fissk to cool somewhat, then add 10 c.c. alcohol through the condenser. Heat to boiling and continue boiling gently for 15 min.

Remove the flask from the condenser and boil the contents until an oily upper layer separates. (See note below.) Add 10 c c. water, cool and pour into a separating funnel. Add 10 c.c. ether (see "Extraction with ether," page 21), shake, allow to stand until two well-defined layers are formed, then run off the lower layer into a beaker and keep this alkahne residue for future treatment. Pour the ether soln. into a dry flask (that used for the hydrolysis will be convenient), connect to a water condenser and partially immerse the flask in hot water contained in a beaker. When the other has distilled off, add 5 drops of the residue to 2 c.c. dichromate mixture

and warm slightly. A strong, bitter-almond odour of benzaldehyde indicates that the residue is benzyl alcohol.

NOTE. If during the boiling down, solid separates and causes "bumping" (as will happen if the ester is benzyl cinnamate) add 10 c.c. water, continue boiling until solid again separates, add 20 c.c. water, heat until all the solid has dissolved, then cool. Extract with 10 e.c. ether, etc.

Identification of the acid constituent of the ester.

. -

Refer to List B of m p.s of benzyl esters (page 110), and proceed as follows:-

If the mp. of O.S. is 80° or near, acidify 2 c.c. of the alkaline residue with glacial scetic acid, heat to boiling and add 2 or 3 drops aq CaCls. A white ppt. indicates an oxalate. If the mp. of O.S. is 42° or below, apply Test (b) under "Thithalio

anhydride" (page 76), using R.G. of O.S.

A vellow-creen fluorescence indicates that O.S. is a phthalate

A yellow green fluorescence indicates that O.S. is a phthalate or succinate.

To distinguish apply Test (e) under "Pathalic anhydride" (page 16), using R.O. of O.S. If the red olour of phenolophithalicias lobation O.S. is a phthalate, otherwise associate. If no yellow-green floorescence is obtained, achilify the alkaline residue with a mixture of equal volume of conc. INO, and water. Filter off the pytel. organic acid, was it with water and dry. Apply the test under "Chinamic acid" (resp. 78).

B. In the flask of a reflux apparatus place 5 g of solid KOH 25 e.e. of alsobel and 2 g of ever. Heat the centeries of the flask to boiling and continue boiling gently for 15 mm. Add 25 e.e. water and dirid 67 20.55 e.e. of lapus in order to remove most of the already 10 per the readies in the flask into a 100-c. explined; make up to 20 e.e. with water and proceed as indicated under "No already has been detected," page 101). Also refer to host C (quer 111), in order to assembly in the flask into a 100-c principle of 0.5 3 the phenol and acid Lidy to be present and proceed accordingly.

Note.

The above procedure is intended for the hydrolysis of the common phenolic esters which are very stable towards appeared. KOH. It is possible, however, that OS may be one of the less common allyl esters, e.g. ethyl phydroxylemosts, mp. 110°, methyl phydroxylemosts, mp. 111°, methyl phydroxylemosts, mp. 121°, methyl phydroxylemosts.

If, there is, no plend is deterted, the lydrelyne should be repeated, many any KOH in the manner described under

A (page 307).

C. From as union B (alone) and all the equivalent of finlayer of upon dust in a six in order to principle contained Filter of the six after the passage of CO.

The only evers of polytyrine plancie likely to be conveniented are assisted and betterated. Owing to the dark-scaleged additions samply cleaned the test for an assiste with Foll.

C. H. 101 110 will be unsatisfactory, while the pptd, benzole acid will be discribured and difficult to obtain white by crystallisation.

Estisfactory results may be obtained as follows :-

If a ppt, has been obtained on acidifying the alkaline residue with HNO, filter it off, transfer it to the hydrolysis flask and add 30 cc. of 23% H.SO. Connect the flask to the water condenser and dietil until about 10 c.c. of distillate is obtained Pure white benzoic acid will collect in the condenser tube and

may be washed out and filtered off. If no ppt, has been obtained on acidifying I c.c. of the alkaline residue with IINO, acidify the remainder of the alkaline residue with 50% H.50. Pour 25 c.e. of the scidified soln, into the hydrolysis flask, add 25 c.c. of 50% H.SO, connect the flack to the water condenser and distil over 10 cc. of liquid. To the distillate add NH,OH until just alkaline

and boil until neutral. To 2 c.c. of the neutral soln, add an equal volume of aq-FeCl, wine red colour (viewed through the depth of the

liquid) indicating an acctate.

See list D for m.p.s of common esters. List A. Methyl esters; esters of phenols and of polyhydric alcohois

#### NOTE.

Except in the case of glyceryl tribenzoate, the time given refers to the hydrolysis of 2 g. with 25 c.c. of 20% sq. KOH. A considerably shorter time is required when 5 g, solid KOH + 5 cc. water, are used.

M.p.		Aydro	dyora
		. 1	5
36° Methyl cinnamate		. 1	
42° Phonyl salicylate (Salol) .			
48° Methyl tartrate			5
54° Mothyl oxalate			
57° Guaiscol benzoate		. 30	
age Phenyl benzoate		. 20	)
70° A.Naphthyl acctate		. 5	
71° p-Cresyl benzoate		, 45	
73° Ethylene glycol dibenzoate		. 60	***
76° Glyceryl tribenzoate	•	. 30	(with 5 g. KOII + 5 c.o. water)
78° Phenyl carbonate	:	. 10	
		15	with alcoholic

List B. Benzyl esters, Hydrolysed in 15 min, with alcoholic

KOH. See under (b), page 108.

- M.s. 39° Benzyl cinnamate 42° Benzyl phthalate
- 42° Benzyl succinate 80° Benzyl oxalate
- iet C. Phenolic esters. Hydrolysed in 15 mm, with alcoholes KOH. (See ander B. page 109.)
  - My 86° Guaiacol carbonate
    - 95° \$-Naphthyl salicylate (Betol)
  - 107° β-Naphthyl benzoate
  - t D. Esters of polyhydric phenols. Hydrolysed in 15 min. with alcoholic KOH. (See under C, page 109.)
    - Mp.
    - 63° Catechol discetate 84° Catechol dibenzoate
    - 117º Resorcinal dibenzoats
    - 123° Quinel discetate 161° Pyrogallol triscetate
    - 199° Quinol dibenzoste
    - autitative hydrolysis of pure esters.

      be determination of the equivalent weight of an ester is valuable
    - the following purposes:—
      i) To ascertain the actual number of acid radicals in the mole-
    - cule of esters of polyhydric alcohols.
      - Example.

        The equiv. wt. of a certain glyceryl benzoate, determined
      - in the manner described on page 112, was found to be 1357.

        The following are the equivalent weights of the three possible glyceryl benzoates, calculated from the molecular

      - Tribenzoate . . 1347
        Ohviously the ester in question is the tribenzoate.
      - To determine the equivalent weight of the acid in an ester of a monohydric alcohol when the alcohol, but not the acid, has been identified.
      - The equiv. wt. of the ester is first found by the method described on page 112.
      - The equiv. wt. of the acid is obtained from this by subtracting the formula weight of the alkyl radical and adding 1 for the H which the alkyl radical has replaced in the acid

, [0]

EXAMPLE.

The equiv. wt. of an ethyl ester was found to be 804.

Equiv. wt. of seid = 80-4 - 200 (formula wt. of C<sub>2</sub>H<sub>1</sub>-) + 1 = 52-4.

On referring to the list of equivalent weights of common acids (page 80) it will be found that the figure obtained is very close to the equivalent weight of malonio acid. Hence the ester is possibly ethyl malonate.

This is particularly useful as an aid to the identification of acids which are readily soluble in water and for whose sails there are no characteristic reactions.

## Practical details.

Weigh accurately a clean, dry flask of 150 c.c. capacity, having a short wide neck and flat bottom. Add 1-1-5 g. of the ester and weigh again.

Into this flask and also into another similar one, introduce 25 c.c. of approximately N. alcoholic KOH and 3 c.c. of water. (The water is necessary to prevent the separation of K salts, such as K phthalate, which would cause violent bumping)

Fit each flask with a cork and reflux condenser and place on a water bath. Allow the contents of the flasks to boil very goth for j hr. (This time has been found to be sufficient for very flashe cotors, such as amyl benzoate, benzyl benzoate, n-butyl phthalse, givernyl tribuczoate.)

Pour 10 c.c. of water through each condenser, also loosen the claup holding the condenser and detach the latter from the fast, raising it so that the under side of the cork is about 1 in above the mouth of the flask. Wash, by means of a wash bottle, this portion of the inner tube which projects through the cett, so that the wash water falls into the flask. Cool the contents of each flask, and phenolphthalein and turtate the alkali present with N. HCl (N. H.SO. gives a ppt. of K.SO. with N. alcohojio KOII)

Norm.

The blank experiment is necessary in order to determine the acid equivalent of the volume of alcoholic KOH employed, since the latter is liable to alter in strength on heating in a glau resel, owing to a tendency for the alcohol to become oxidized by the air, with subsequent resinification of the alichytle formed, and also owing to a object action on the glaus.

If the ester is one which is readily hydrolysed (for example, in 15 min. by 20% aq. KOH) and yields a water-soluble alcohol, agneous N. NaOH may be employed.

In this case bull the contents of the flask gently over a gause, noting the time taken for the !

then continue boiling for a further period of about \( \frac{1}{2} \) of this time. If the ester is miscible with the NaOH soln, boil for 15 min

If v e.c. = the difference between the two titrations

and w g. = the weight of ester taken

Equiv. wt. of ester =  $\frac{w \times 1,000}{v \times \text{factor of acid}}$ 

B-Ketonic esters

181° Ethyl acetoacetate. CH, CO CH, COO Et.

Fix a 100-cc. flask, having a short wide neck and fits bottom, with a cenk and a reflux condenser. Remove the flask from the condenser, pour into it 25 cc. of 20% at, KOH and 5 cc. of 0.5. and add some porous pot. Connect the flask again to the condenser, heat the contents to boiling and containe boiling for 5 min. Allow the contents of the flask to cool somewhat, then pour 10 cc. water through the condenser. Disconnect the flask and stach it to a sloping condenser. Distil, using a £1. as the receiver, until 10 cc. of flatiblists is obtained. Pour the silkance residue in the ski into a beaker and keep it for the treatment described below. Rimso out the flask with water.

Treatment of the distillate.

(4) To 2 c.c. add an equal volume of ½% ag, sodium introprusside, then add 2 drops aq. NaOH. A wine-red colour, which is changed to violet-red on scidifying with acetic acid, indicates that acetone is one of the products of hydrolysis.

(4) Four the remainder of the distillate into the empty flask which was used for the hydrolysis and add 20 c.c. dichromate mixture and some porous pot. Connect the flask to the sloping condenser and datill, using a t.t. as the receiver, until 5 c.c. of distillates is obtained. To 2 c.c. of the dillates add an equal volume of 20% as, ROH, heat to boiling, and continue boiling for i min. A pitch which, followed by a yellow pyt., changing coverage, with a disagreedule olour (due to the formation of acetaldehyde rein) indicates that ethyl alcohol is one of the products of hydrolynia.

Treatment of the alkaline residue in the beaker.

Acidify with 50% H<sub>2</sub>SO<sub>4</sub>; a vigorous effervescence indicates the presence of K<sub>1</sub>CO<sub>4</sub>.

c, n. to:

Ц preparation from it of adiabatement,

mp por as follows --Forme I e. of OS. Into a der 100 ac. first and sell a mirture of 3 ea. ea. of enor. HNO, and cone. H,50, Heat to builing and continue builing for i min. Cred, pone into 60 e.e. water, cool and shake Filter, wash the solid with cold water, crystallien twice from alcohol, dry, and determine the m.p.

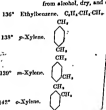
(2) cycloffexano is unaffected by the treat ment.

110° Toloros. C.H. CH.

111

Ph

Preparation of 2: 4 dinstrotoluene, mp. 70°:-Pour 1 c c. of OS. into a dry 100 c.c. flask and add quickly a mixture of 3 c.c. each of fuming HNO, and cone, H.SO. (The reaction is violent.) Cool, pour into 50 c.c. water; cool and shake. Filter, wash the solid with cold water, crystallise twice from alcohol, dry, and determine the m p.



Oxidation test:-Into a 150 c.c. wide-mouthed flask pour I c.c. of O.S. and add 5 g. solid KMnO4, 80 c.c. water, 10 drops aq. NaOH and two or three pieces of porous pot. Fit the flask with a reflux condenser, heat the contents to boiling and continue boiling fairly rapidly for 2 hr. Cool, and pass SO, until any purple colour and the brown ppt. have disappeared.

If (a) a clear soln, is obtained, filter from unchanged hydrocarbon, cool well and shake vigorously. Filter off the solid which crystallises out, wash it carefully with cold water and dry. Apply tests for phthalic acid from 771.

> Positive results indicate that O.S. is o-xylene. Confirm the identity of O.S. by preparing o-xylenesulphonamide (see below).

If negative results are obtained, determine the m.p. of the solid. A m.p. of 121° (or near) indicates that the exidation product is benzolo seid and that O.S. is ethylbenzene

(b) a bulky white ppt. is obtained, filter, wash the solid with water and dry. Apply the test given under isophthalic and terephthalic acids (page 78), and distinguish between the acids by preparing the dimethyl ester.

Identification of the solid as (1) isophthalic soid, indicates that O.S. is m-xylene.
(2) terephthalic soid, indicates that

O.S. is p-xylene.

Preparation of p-xylenesulphonamide, m p. 114.

To 1 e.e. of the hydrocarbon in a dry t.t. add 2 e e. cone. II SO., immerse the end of the t.t. in boiling water and shake until the hydrocarbon has completely dissolved. (1-1 min.) Cool and pour into 20 e.c. of a saturated soln, of common sait. Cool. and shake or stir vigorously. Filter off the ppt, of sodium aylenesulphonate and wash it with saturated salt soln. Dry the ppt, thoroughly in a steam oven. Place the dried salt in a porcelain dish, add 4 g. PCl, and grind the two substances intimately together with a peatle. Heat on a rapidly boiling water bath for 10 min. stirring the mixture periodically. Ocal the mixture, add 10 c.c. cold water and stir. Wash the contents of the dish into a separating funnel and extract with ether (see page 21). Dutil off the other, thormoraly mix the reside of sulphonyl chloride wall

C, II, [0] 118

2 g. powdered ammonium carbonate, and heat on a rapidly boiling water bath for 15 min. Cool, B.D. add 20 c.c. cold water, stir well and filter. Wash the sulphonamide with water, crystallise it from alcohol, dry, and determine the m.p.

154° Anisole. (Phenyl methyl ether.) C.H. O.CH.

Preparation of dinitroanisole, m.p. 88°:-Dissolve 1 c.c. of O.S. in 5 c.c. conc. H.SO.; col. Add I c.c. cone. HNO, in portions (about 5 drops at a time) cooling after each addition. Pour into 20 c.c. water, cool and shake. Filter, wash the solid with cold water, crystallise from a small quantity of alcohol, dry, and determine the mp

O-Ma 171° o-Cresyl methyl ether. 172° Phonetole. (Phenyl ethyl ether.) C.H. O.Et. O-Ma 175° p-Cresyl methyl other. ČH(CH.). 177° m. Crosyl methyl ether.

To I e.e. of OS, sold I ee, of cone. If SO,; gently shako.

If Q B.

A. does not dissolve completely in the some. II,SO, pour I ce. of OS, into a 100 ce. widemouthed flack, add 30 co. dichromate misture and two or three pieces of parinis jas. Fit the flack with a reflux numberser, heat the contents of the flash to boiling and continue beiling fairly armily for 3 to Alt to) en water and filter

Wash the solid with water until free from Cr compounds and dry. Apply the test for terephthalic acid (page 78) A positive result indicates that O.S. is p-cymene.

B. Dissolves completely in the conc. H.SO. [soln. may be yellow, pink, or light red brown in colour), add another 9 c.c. of conc. H.SO, and cool. Add 1 c.c. conc. HNO, in portions [about 5 drops at a time), cooling after each addition. (Soln. will become reddash or greenish brown.) Four into 60 c.c. water, cool and abade.

If (a) a pale yellow conclains a obtained, yielding a pale yellow colid on shaking, filter. Wash the solid well with cold water, crystalline from alcohol, dry, and determine the min.

Mp of nitro derivative :67° indicates that O.S. is o-creayl methyl

86" indicates that O.S. is phenetole 91" indicates that O.S. is m-creayl methyl other

(b) no yellow emulson is formed, but the matture froths on pouring into water, yelding an orange-brown ando, with hitse or no solid, proceed with the oxidation text described under "B. p. 130"-142" (page 116), using 2\cdot g. solid KiMoQ. do c. water, 5 drops a NoOll and I.c. of OS. The KiMoQ will sustally be completely reduced in It he.

Identification of the oxidation product as anise acid, mp. 184°, indicates that O.S. is p-cress! methyl ether.

CH: CH CH.

222° Anethole,

mp 21°. Olour of anseed,

in) To jun, layer in a ts. of a soin of fir in OCI, all I drops of O.S.; state,
--brown return rapidly disappears, due to the

2.0 ho develous at as food oldself a la succession of 0.5

ſΙ

(3) Oribition to anine acid, mp. 184°. lata a Sinee, flack pour 50 e.c. dichromate printers and I e.e. of O.S. Heat, shaking road fragmently, until a vigorous reaction commences then remove the fisme. When the rignous to action has crased, cool, filter off the solid, and wash is with water to remove green Creality & the solid to 20 c.c. boiling dil. HCl, then add fow crystale of Na SO, boil and stir for 5 m Add 10 ce. cone. NII,OII, boil for 5 min, th cool and filter. Acidify the filtrate with on HCl. filter off the solid, wash it with with crystallise from dilute alcohol, dry, and determine

the m.p. O.S. sinks in water. See notes under "O.S. floats on water" (page 115). O-Mo

0-3fe B.D. 205° Catechol dimethyl ether (Veratrole)

214° Resorcinol dimethyl ether

Preparation of dibromo derivative. (The bromination should be carried out in a fume cupboard.) Into a 100-c.c. flask pour 1 c.c. of O.S. and add 1 c.c. bromine. When the vigorous reaction has ceased, add 20 c.c. aq. NaOH and shake. Dilute with water, filter, wash the solid with cold water, crystallise from alcohol, dry, and determine the

m p. (If the product possesses a brown colour, recrystallise until it becomes white.)

M.p. 91° indicates dibromo catechol dimethyl other. resorcinol .. 141°

ETHERS AND HYDROCARBONS 121

B.p. To 1-in, layer in a t.t. of a soln of Br in CCl, add

3 drops of O.S. and shake.

-brown colour rapidly disappears, due to the presence of a double bond in the molecule of O.S. Alkalina KMnO. - representite acid, in p. 223°.

O-Me

263° a.Naphthyl methyl ether.

To 2 cc. of a cold saturated soln of pictic acid in benzene add 2 drops of O.S.,

-red crystals of a picrate form methoxyl radical = 196

O.S. solla.

"Twodure for the alentification of O.S. (Rec Note 2, page 113.)

Determine the mp, then refer to the list of mps of hydrocarbons and ethers. If one of these mps is identical with, or near to, that of O.S. apply any tests given, and prepare the detrivative there indicated. (See sections on "Crystallisation"

and "Drying of substances," pages 10 21)
In the case of others, where no derivative is indicated, determine
the b p. and also the percentage of methods) or ethods) radical

the h p, and also the percentage of methoxyl or ethoxyl rad (For method of determination, see page 120)

Mp
21° Anethole Crystalline mass Olour of amoved Bp 222°
See "Anethole" uteler OS fluts on water"

(page 119)
26 Diplomylmethase C.H.CH.CH. Crystalline mass.

Olour rewriting cranpes. Bp 2012 Oxidation to beneghtmore, Call, CO Call, identified

by contrast on the place of the

alcohol, filter and wash the sold of his Ferresilies the wild two from Model of all determine the or p.

phenyl ather (Diphenyl cusls) CH,OCH, to talline mass. Germine the other 8.2 of Frepretion of 4 f disable deviation, E. Pti. Contraction should be carried out in time spot as the reaction is violent and cropins final

3

37° A.Naphthyl ethyl ether. O'Et Powerful &co cdocs.

B.p. 282°.
Picrate, m.p. 104° (see page 143).
°/ ethoxyl radical = 26°1.
OMe

47° Pyrogallol trimethyl ether, OMe

87 methoxyl radical = 55-4.

52° Diberrit CH<sub>3</sub>CH<sub>2</sub>CH<sub>3</sub>CH<sub>4</sub> B<sub>4</sub> B<sub>5</sub>

Orderion to fearne and, mp. 121'
In a 100-ce, wide months flash place 2 if Millog, 40 ce, wider, 5 ctype 44, 2003, 14 if Nillog, 40 ce, water, 5 ctype 44, 2003, 14 if the Nillog, 40 ce, water, 5 ctype 44, 2003, 14 if the Nillog, 40 ce, water, 5 ctype 44, 2003, 14 if the Nillog 40 ce, water, 5 ctype 44, 2003, 14 if the Nillog 40 ce, water, 5 ctype 44, 2003, 14 if the Nillog 40 ce, water, 5 ctype 44, 2003, 14 if the Nillog 40 ce, water 10 ce, water 1

liquid to believe the disappeare, and liquid to believe there from mediand letearbon, cool word and shall be with old wife. Filter of the solid and wash is with old wife crystalline from water, they and determine its #? M.p.

O-Me

55° Quinol dimethyl ether.

% methoxyl radical = 44-9.

radical  $\approx 44.9$ .

70° Diphenyl. C.H.-C.H. B.p. 254°.

Preparation of ppf differentiative, mp. 168\*—
In 100.co. beaker place ; s. 0.05. and an amount of iodine roughly equal in bulk to a half res. Add a soln. of I co. Br in 4 co. CCI, Allow to stand 30 min, then evaporate for dryness on a water bath. Add 10 cc. boiling aq NaOII and str. in order to remove as much iodine as possible. Dutto with 10 cc. water, fifter and wash the solid with water. Crystallies the solid from aestons,

dry, and determine the mp.

72° \$\beta\$-Naphthyl methyl ether (Nerolin.)

 $\bigcap^{n}$ 

Powerful odour of orange blossom. B.p. 274\*. Picrate, m.p. 118\* (see page 143)

methoxyl radical = 19-5.

80° Naphthalene. Characteristic odour B p 218°.

Preparation of picrate (yellow), m.p. 149\* --

Dissolve 4-1-in. Layer of powdered O.S. In a t. i. in l ce, cold benzere. Add 2 ce, of a saturated soln, of picrio acid in hemzere and rhake. Filter off the ppt, and carfelly wash it with a few drops of cold benzere. Press the ppt, between three paper, dry over a small flame, and determine the mr.

92° Triphenylmethane. (C.H.),CH.

In a dry tt. place D | g of O S. and add 2 c., furning INO; a low to stead S min. Nearly Chief to take with rold water and shake There of the 17th tention composed and wash it with water. Transfer the pre, to a t.t. and damobre it in 10 c.c. placid article acid. Add Zn dust wath the deep red colour, which is formed as fact, dangresse, 124 C, H, [O]

M.p.

then filter. To 2 c.c. of the yellow filtrate 2 R.G. of PbO, and shake,
—intense red colour, due to the formation
pararosaniline.



95\* Acenaphthene.

Picrate (orange), m.p. 161°. (For preparation and under "Naphthalene," page 123.)

100° Phenanthrene.



Picrato (yellow), m.p. 143°. (For preparation secondar "Naphthaleno." After mixing the solns, heat just to boiling, cool and shake)

113° Flaoreno.

Oxidation to the ketons, fluorenoite (yellow), khatched by conversion to its phenythydrazone, m p 18. In a 100-c. ox while-monthed flask place 1 g. of (19. 3), swimm dichromate and 5 c.c. glacial series should be contents of the flask to bolling and contents of the flask to bolling and contents losling gently, with previous shading round, \$\xi\$

I be.
Add through the condenses 50 a.s. of beling wife
Transfer the contents of the flack to a (6) a.
distilling flack, make up to volume to abed
20 to e, with water and add 2 or 3 purces of procepest. Connect the flack to a condense and led
the contents represently. Filter off the yellow
floor notes which passes over with the steam, and
perpart its phroylly-frazone in the manus the

ertied under "138 mild," (page 69) Crystallion the phenythydrames from absolut (the derivative in sparingly milded in absolut, whereas it is correct in smally milded dry, and determine the mile. M.D.

125° Stilbene, C.H. CH:CH C.H.

(a) To t-in, layer of O S, in a t.t. add 5 c.c. dichromate mixture, heat to boiling and continue boiling for min.,

-bitter almond odour of benzaldehyde.

(b) Pour 5 c.e. Br water into each of two t.t.s. one add the equivalent of 1-in. layer of OS in a t.t. Stand both tubes in boiling water for 1 mm . then take them out and shake vicerously. It will be found that the liquid in the tube containing the hydrocarbon has become colourless fowing to the presence of a double bond in the molecule of the substance), whereas in the blank test the colour of the bromine persists

CH

Anthracon

(a) Add 2 R G, of O S, to 2 c.e. of a saturated soln, of pierie acid in benzene and shake. -deep red soln., due to the formation of a red

picasto. m

(b) Oxidation to Anthropyin

Weigh out in a 100-c c, wide-mouthed flask 1 g. of CrO, and add a mixture of 1 e.e. water and D c c. glacial acctic acid. Heat until the OrO. has descired: cool.

Add I g. of O.S., fit the flack with a reflex condenser and heat the contents to buling | [1] the mixture froths up, remove the flame for a moment ) Continue boiling gratly for 10 min. shaking the flask round periodically. Allow to ered somewhat, then add 50 c.c. water, Filter, and wash the wish with water until it is a clean Stillow colour. Apply the fillraing feet for anthraminone ....

In a t Citier 2 RG of the yellow wild and an equal bulk of Zn dust Add 5 c.c sq NaOH, heat to boiling and continue boiling for | 22. -deep red colour.

Filter while hot into a t.t. and shale, -red colour rapidly disappears, owing to exists by the air, and a flocculent pale relice pet anthraquinone separates. The phenomens cribed may be repeated by adding Za dut si boiling, etc. The red colour is due to the Na



DETERMINATION OF THE PERCENTAGE OF METHOXIL OR EVI RADICAL IN METHYL OR ETHYL ETHERS

Perkin's modification of Zeisel's method.

The method depends on the fact that when a compound contains methoxyl or ethoxyl groups is heated with hydrixis the methyl or ethyl groups are converted into methyl or iodide, which when passed into an aqueous alcoholic scin. of a nitrate, is decomposed with the formation of silver iodide. the weight of silver iodide produced the percentage of metho or ethoxyl radical is calculated. Reagents employed.

Hydrodic send. The constant boiling mixture is required, 127° and d. 1-70, containing 57% of HI. Buch an acid, for in Zouel estimations, may be purchased

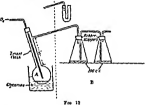
A punes alcoholic piler nurate polutum Disselva 2 g Ag 10. See. distilled water and seld 43 ag. absolute alcohol. The sel must be kept in the dark, and if not clear, the required quantil should be filtered through a dry filter pares into the absorption Cask .

Carbon durate This may be obtained from a Kipp's apparaisa the gas being washed from from traces of HCT and any HA (origin from impuritors in the marking by passing through a dilute site of \$270 and then dred by passing through some Highly

A roundus

The apparatus used in illustrated as Fig. 12. The being north is the special Zound flock A wite as my air manderione must bear been the tribulational and and any notion promote it amount he model that the sort of the concentral train my the first flack frombonised 2. a - ALTIS, mile | terminates both steres the section of

e liquid, while that in the second flack (containing 15 c.c. of (NO<sub>a</sub> soln.) dips well below the liquid.



refere:—

emore the cerk and tube from A, and by means of a thirde

wh, the end of which reaches well into the both, introduce

c. of the hydricule acid into the fask. Allow the stem of
funnel to drain and then exercitly remove the funnel so that

roys of acid are left on the neck of the fask. All as few pieces

arous not to the content of the fask. here relates the cerk

tube so that the end of the tube is just above the acid, tech to the side-tube of the dutalling flack a U tube containing 'e.e. of the AgNO, soin. Baue the temperature of the plyerene to 100° and pass above current of CO, through the apparatus the issuing gas does not give a turbelity with a quantity of AgNO, soin. Cities to remove any voisited impurites from ydriche and.) Allow the apparatus to cool, discenser; the he and replace it by the absorption apparatus B.

igh out accurately in a small plan take about 0.3 g of the person the occh sarrying the CO, take and take the take inling the other down into the seal of necessary peak in down the CO, take). Furthy replace the cost and CO, take, pancurrent of CO, through the apparatus (about I helding purrouncest of CO, through the apparatus (about I helding purlatives; the II SO, would bettle, and heat the previous to 140°. A whate deposit to compound of salver action and is seen beginn to form out the service of the hight in the sale and productly settles to the bettern, but analy only a sequence in second faul.

128 After about 20 min. raise the temperature of the glycerine and the hydriodic acid boils gently, but not so vigorously as to care distillation of the acid into the side-tube of the flask. Octaor heating for 40 min, then disconnect the flasks and substitute for them the cleaned U-tube, containing a few c.c. of the AgNO, xh

Heat for a further period of 20 min., and if no appreciable quantit of ppt forms, the distillation of the methyl or ethyl todale may be considered complete; otherwise pour the soln into one of the flasks, place a fresh quantity in the U-tube, and repeat the process

until the formation of ppt. ceases.

To 50 c.c. distilled water contained in a 250-c.c. beater all 10 c.c. dil. HNO, and heat to boiling. Gradually add the content of both flasks and rinse out the flasks, the connecting tube sal the U-tube with hot distilled water into the beaker. Continue boiling for some time in order to drive off the alcohol, and to decompose completely the white, somewhat stable, double sait of silver iodide and silver nitrate which has been formed by the interaction of the methyl or ethyl iodide and the alcoholic AgNO.

Add boiling distilled water, if necessary, to prevent the volume soin. becoming less than about 50 c.c. and break up any lumps of Fit. by means of a glass rod. Allow the soln, to stand in the dark of an hour, then filter off the ppt. of AgI into a tared Gooch or sintered

glass crucible, wash, dry at 100°, and weigh.

234 8 g. of AgI correspond to 31-02 g. of O-CH, or 4504 g of

EXAMPLE :- 0-3150 g. of a methyl ether gave 0-1630 g Agl % methoxyl radical = 31 03 × 0 488 × 100 = 19 6. The results obtained with athory compounds are generally somewhat her.

ADDITIONAL COMPOUNDS CONTRIBUTED C and II, on C, II, AND O

(1) Consturated compounds may be detected by the fidicaled tret .--

To I in layer in a \$8, of a soln, of Be in \$17, att one se two der pa of O.S. and shake wherein echoir rapully desapposes Prose the ecotroits showly out of the title, me coppers grafitue of lille fience, including that will turn and me mitatibution had

(2) Fix further perspecture and descentives a turn manipularies with abunda has measured

to The shore of the estations if now a hydrometric, is studied

Liquide. Float on water. B.p.

30° ico-Pentane. (CHa), CH-CHa-CHa. Petrol-like odour. 35° Diethyl ether. (C.H.),O. 36° n-Pentane. CH. [CH.] CH. Petrol-like odour,

(Standard pentane, boiling range 25°-40°.) e9a n-Hexane. CH. [CH.]. CH. Petrol-like odour. 69° Di-isopropyl ether. (CH,),CH O CH(CH,), 900

Di-n-propyl ether. (CH, CH, CH, CH,),O. 98\* n-Heptane. CH, [CH,] CH, Petrol-like odour.

125° n-Octane, CH, [CH,], CH, Petrol-like odour. 146° Styrene. C.H. CH.CH. Unsaturated.

155° Pinene. C<sub>18</sub>H<sub>18</sub> Odour of turpentine. Unsaturated.

164° Mesitylene. C.H.(CH.), (1.3:5). 176° Cincolo, C<sub>10</sub>H<sub>14</sub>O. Odour of eucalyptus. (Ether.)

176 Limonene, CraHra. Odour of lemon, Unsaturated. Indene. C.H. Unsaturated.

192° Decahydronaphthalene. C.H. 206° Tetrahydronaphthalene, C., H.,

Sink in water.

240° α-Methylnaphthalene. C, H, CH, Picrate, m.p. 141°. 295° Dibenzyl ether. (C.H. CH.) O.

Solide Mp.

32° β-Methylnaphthalene. C<sub>1,H1</sub>·CH<sub>2</sub>. Picrate, m.p. 115°. Cinnamyl alcohol. C.H. CH = CH CH OH. See page 38. 42° I-Menthol. CasH 110H. Odour of peppermint. (Saturated

48° Cetyl alcohol. CH [CH ] CH OH. 51\* l.Camphene. C.H., Unsaturated. 68°

Diphenylcarbinol. (C.H.,),CH-OH. (Secondary alcohol.) 116° Terpin hydrate. C,H,O, H,O.

162° Triphenylcarbinot. (C.H.), COH. (Tertiary alcohol.) 203° d-Borneol, C10H1,OH, Odour like camphor, (Secondary

F.

C. II. [0] 128

After about 20 min. raise the temperature of the glyceme and the hydricdic acid boils gently, but not so vigorously as to care distillation of the acid into the side-tube of the flask. Content heating for 40 min, then disconnect the flasks and substitute for them the cleaned U-tube, containing a few c.c. of the AgNO, sola

Heat for a further period of 20 min., and if no appreciable quantity of ppt. forms, the distillation of the methyl or ethyl fodde my be considered complete; otherwise pour the sola, into one of the flasks, place a fresh quantity in the U-tube, and repeat the process

until the formation of ppt. ceases.

To 50 c.c. distilled water contained in a 250-c.c. bester add 10 c.c. dil. HNO, and heat to boiling. Gradually add the contests of both flasks and rinse out the flasks, the connecting tube and the U-tube with hot distilled water into the beaker. Continue boiling for some time in order to drive off the alcohol, and to decompose completely the white, somewhat stable, double salt of silver iodide and silver nitrate which has been formed by the interaction of the methyl or ethyl iodide and the alcoholic AgNO,

Add boiling distilled water, if necessary, to prevent the rolume becoming less than about 50 c.c. and break up any lumps of pp. by means of a glass rod. Allow the soln, to stand in the dark of an hour, then filter off the ppt, of AgI into a tared Gooch or sintered glass crucible, wash, dry at 100°, and weigh. 234-8 g. of AgI correspond to 31-02 g. of C

O-C.H.

EXAMPLE :- 0.3150 g. of a methyl other :

% methoxyl radical =  $31.02 \times 0.468 \times 100$ obtained with ethoxy compounds are g

ADDITIONAL COMPOUNDS CONTAINING AND O

Notes:

(1) Unsaturated compounds may be d test :--

To 1-in. layer in a t.t. of a two drops of O.S. and shake-Pour the contents slowly out of ... of HBr fumes, indicating th

(2) For further properties and deri work should be consulted.

(3) The class of the substance, if

## SCHEME II

## Compounds containing CI, Br, or I.

Follow the appropriate procedure either under "O.S liquid" (below) or under "O.S. solid " (page 132).

## O.S. liquid.

Apply the following tests in the order given.

(1) To 3 c.e. of distilled water in a t.t. add one drop of O.S., heat to boiling, continue boiling with shaking for 1 min., then cool. To the clear soln, (if necessary obtained by filtration) add 2 e.c. of dil, HNO, and I c.c. of an AgNO. If a definite ppt. of Ag halide is obtained (ignore a slight milkiness) see "Acid

halides, etc." (page 150), otherwise apply Test 2. (2) To 1-in. layer of 2 : 4-dinitrophenylbydraune in a dry t.t. add

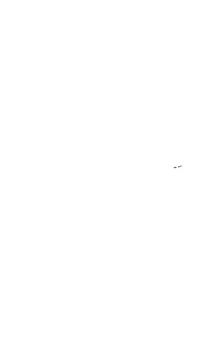
2) c.c. of alcohol, then add | c.c. of cone 11,50. Warm and shake in order to dissolve all the solid. To the soln, add & c.e. of O.S., shake and allow to stand for 5 mm unless a ppt. forms in a shorter period of time. Finally cool and shake the contents of the t.t. If a ppt. is obtained, retain it and see " Halogen. aldehydes and halogen ketones" (page 138), or if there is no

ppt, apr.ly Test 3. (It is necessary to apply this test before Test 3 as chlorobenzaklehydes dussolve somewhat in aq. NaOli and are re-

precipitated from the alkaline soin, by HCI) (3) To 5 drops of O.S. in a t t. add 3 e.e. of approx. 2N. NaOH close the mouth of the tube and shake vigorously for I min If any O.S. has remained undissolved add 3 c e. of water, ther shake and filter the contents of the 11. Acids the soln, o filtrate with come. HCL. If a white emulsion or an oil is ob tained, see "Habsen-phenols" (page 136), otherwise apply Trot 4. (No account is taken here of halogen enlatituted highe

aliphatic acada, which, if sparingly soluble in water, would give a postive result in this test.) (4) To 2 c.c. of alrebol add one drop of O.S. and 10 drops of approx

 $\frac{N}{2}$ NeOH, shake round and then all a deep of phenotylethalou



colourless see "Aliphatic halogen-acids" (page 133) or if a red colour is produced apply Test 2.

- (2) To \(\frac{1}{2}\) in layer of O.S. in s t.t. add 2 c.o. of approx. 2N. NoOH, heat to boiling, continue boiling gently for 1 min., then cool. Add s drep of phenolphthalein soln., then add dd. HCl until the soln, just becomes colouriess. Now add 1 c.o. of aq. HgGl, heat to boiling, continue boiling for 1 min., then cool and shake, by A white pp. of HgCl, due to the reduction of the HgCl, by a formate produced by hydrolysis) suggests that O.S. 10 a hydrate or alcoholate of a landogm-aldebyle. See page 13 a.
- (3) To 4 in. layer of O.S. in a dry tt. add 3 c.c. of alcohol, heat just to boling, then cool and sake; filter if any solid is press. Using this alcoholic soln, instead of the § c.o. of O.S. proceed as in Test 2 (page 131). If a ppt is obtained retain it, and son "Halogen-alchydre and halogen-lectones" (page 138), otherwise apply Test 4.
- (4) To j-im layer of O.S. in a t.t. add 3 c.c. of approx. 2N. NaOH, close the mouth of the tube and shake vagorously for 1 min. If any O.S. has remained undansolved add 3 c.c. of water, then shake and filter the contents of the t.t. Acidify the sols, or filtrate with come. HCl. If a ppt. or a whote emulsion is obtained, see "Aromatic Indogen.acids and Indegen.phenole" (1945-138), otherwise apply Test 5 unless O.S. has a camphor-
- lke odour, in which case see first "Chlorbutel" (page 138).

  (5) Proceed as in Test 6 (page 132) using the equivalent of 1-in.

  layer in a t.t. of O.S.

## Metal present.

Only alkali salts of halogen-acids are here considered.

Procedure:—

Prepare about 5 c.c. of a cold, concentrated soln. of O.S. and acidify with cone. HCL.

- If there is obtained

  (a) no ppt, proceed as indicated under. "Aliphatic halogenacids" (page 133).
  - (6) a ppt., filter it off, wash it with cold water, dry, and proceed as indicated under "Aromatic halogen-acids" (page 135).

## HALOGEN-CARBOXYLIC ACIDS

## Allphatic balogen-acids.

The halogen-substituted acetic acids included in this subsection possess a sharp odour, are deliquescent if solid, and have a blistering action on the skin. soln. If the soln, remains colourless see "Aliphatic halogen

acids" (page 133) or if a red colour is produced apply Test 5 (5) To 1 c.c. of acetyl chloride in a dry t.t. add about i c.c.d O.S. If within 2 min. a vigorous reaction occurs (i.e. bubble are freely evolved) with evolution of HCl fumes, see "Halogen alcohols" (page 137), otherwise apply Test 6.

(6) In a 100-c.c. wide-mouthed flask place 2 g. of solid KOH and 10 c.c. of alcohol and heat under a reflux condenser until all the KOH has dissolved. Pour 2 c.c. of the soln into s th for use as a blank test. To the remaining soln, in the first add i c.c. of O.S. and heat to boiling under the condenses. If a considerable ppt, separates, no further boiling is necessif. otherwise continue boiling for 10 min. Cool, acidify with al HNO, cool well and shake, then filter. Also acidify the 2 ce. of alcoholic KOH in the t.t. with dil. HNO. To 2 cc. d each acidified soln, add I c.c. of aq. AgNO,

(a) a ppt. of Ag halide greater than that obtained in the black test see "Aliphatic halogen-hydrocarbons, etc." (page 140)-(b) no greater ppt. of Ag halide than that obtained in the blank test proceed as under (b), page 99, in order to seet.

tain if O.S. is an ester of an aromatic halogen seid. If O.S. is not an ester, see "Aromatic halogen-hydrocarbons" (page 146).

O.S. solida Follow the appropriate procedure either on page 133 or below, according to whether a metal is present or not.

No metal present. To a measured 1-in, layer of powdered OS. in a dry 1-in. 11. add 5 c.c. of distilled water, heat to boiling with shaking, then cool well and shake. If O.S is completely in solution, apply Test 1 (below) or if it is not completely in solution proceed as under (1). page 131, using R.G. of OS. If no ppt, of Ag halide is obtained, apply Test 3 (below).

(1) To 2 c.c. of the aq soln. of OS add a drop of an FeCl, Il a blue, violet, or green colour is observed see "Halogenphenols" (page 136); if none of these colours is obtained then to 2 c c. of dil. NH OH add R G. of OH, and shake for 1 min If a deep brown or deep red colour is observed see " Halogenphenols" (page 136), otherwise to a soln, of It O of Og in

2 ee, of water add 5 drops of approx. No NaOII, shake and then aid a drop of phenolphthalein soin. If the soin remains

and cool until solid separates. Filter, wash the solid with cold water, crystallise it from water, dry, and determine the m p.

Bromo-acids

M.p. 50° Bromoscetic acid. CH, Br COOH. Follow the procedure under "Chloroscetic acid" (page 134) when similar results will be obtained. Aromatic halogen-acids and halogen-phenols.

Determine the m p. of O.S. (if below 100° see "Halogenphenols," (page 136), and refer to the appropriate list of m.p.s of acids. Method 1 (page 81) should be used for the determination of the equivalent weight. For derivatives see page 80.

Chloro acide. M.P. COOH	Equit. to
140" o-Chlorobenzoic acid. Anfide, m.p. 142". p-Nitrobenzyl ester, m.p. 106".	
153° m-Chlorobenzoic acid. Amide, m.p. 134°. p-Nitrobenzyl ester, m.p. 110°. COOH	156-5
230° p-Chlorobenzoic acid. Amide, m.p. 179°. p-Nitrobenzyl ester, m.p. 129°.	
decomp. Tetrachlorophthalic acid. p.Nitrobenzyl ester, m.p. 180°.	151-9
M.p.	Equip. w
147° o.Bromobenzoie acid.  Amide, mp. 157°.  PNitrobenzyl ester.  mp. 110°.  155° m.Bromobenzoie acid.  COOH  Amide, mp. 155°.  p.Nitrobenzyl ester.  mp. 105°.  Dr.  Dr.  Dr.  Dr.  Dr.  Dr.  Dr.  D	200-9
195° Dibromorinnamic acid. C.H. CHBr CHBr C	OOH 307.9

O.S. Hawld.

Chammente. Apply the following test for " Dishloroscetic soid "CHCI, COOK h p. 100" Into a small flask pour 1 ee, of O.S. and 10 ee d and KOH, heat to beiling and continue beiling gratly in 5 min (Sola becomes yellow) Acidify 2 cc. of the sola with glarial acetic acid, bent in builing and add 2-3 drops ac Cally -immediate white ppt. (Ca ovalate, due to the formation of

osslic sekti Confirm the elentity of O.S. by preparation of the aniline salt, in p. 122 as follows :- Mrz intimately in a porcelain dish I ca each of OS and freshly distilled aniline. When the mixture has become solid, triturate with 3 cc. ether. Filter, wash the adcarefully with ether until white, dry, and determine the mp.

O.S. solid.

Chloro-neide Proceed as indicated under "Trichloroscetic acid."

If negative results are obtained follow the procedure under "Chloroacetic acid."

55° Trichloroacetic scid. CCI, COOH.

Apply the tests under "Chloroform" (page 140) using R.G. of O.S., when similar results will be obtained owing to chloroform being first formed.

63° Chloroscetic acid. CH,Cl COOH. (a) In a boiling tube place the equivalent of 1 in larer of O.S. in a t.t. and add 10 c.c. aq. NaOH. Heat to boiling and continue boiling for I min. Add the equivalent of j-in. layer in a t.t. of solid KMaO. and shake until the green colour has disappeared, then filter. To 2 c.c. of the filtrate add I c.c. glacial acetic acid, heat to boiling and add 2-3 drops

-immediate white ppt. (Ca oxalate, due to oxalic acid formed by exidation of the glycollic scid produced by hydrolysis).

(b) Preparation of phenoxyacetic acid,

с.н.о сп. соон. m p. 96°:-

In a 100-c c. wide-mouthed flask place I g. of O.S., 1 g. of phenol, 11 g. KOH and 10 c.c. water. Fit the flask with a cork and reflux condenses, heat the contents to boiling and continue boiling for 5 min. Cool, acidify with cone. HCl. shake

67° sym-Trichlorophenol,



Benzoate, m.p. 70°, Methyl ether, m.p. 60°.

93° sym-Tribromophenol.

Acetate, mp 82° (crystallise from alcohol). Methyl ether, m.p. 87°.

## HALOGEN-ALCOHOLS

#### O.S. liquid containing Cl. Procedure for identification.

Determine the b.p. and refer to the following list of b.p.s. if one of these b.p.s is identical with, or near to, that of O.S., confirm the identity of O.S. by applying to it any tests given, and by proparing and determining the m.p. of the derivative there indicated.

## B.p.

127° Ethylene chlorhydrin, CH,OH-CH,CL Miscible with water.

- (e) Pour into a bolling tube ê c.c of O.S. and 10 c.c. aq NaOil, heat to boiling and continue boiling for 1 min. Add an amount of solid KMnO, equivalent to 1-ln, layer in a t.t. and shake until the green colour disappears. Filter, actifity 2 c.c. of the filtrate with glacial acetic acid, hat to boiling and add 2-3 drops of aq. CcU<sub>2</sub>. —white pp. (Ca casalac, due to oratic acid having been formed by the oxidation of the ethories of the colour disappears.
- (b) Preparation of allylene plycol dibenzode, m.p. 73°,— Into a 100-c.c. conical flask pour 2 c.e. of O.S. and 20 c.e. aq. NaOII, heat to boiling and continue boiling gently for 5 min. Cool, add 2 c.e. benzoyl chloride and 20 c.e. aq. NaOII. Oork the flask and shake until solid separates, then shake for a

HALOGEN, C, H, [O]
M.p. COOH

251° p-Bromobenzoic acid. Amide, m.p. 189°. p-Nitrobenzyl ester, m.p. 139°.

# HALOGEN-PHENOLS

Procedure for the identification of O.S.—

Determine the bp. if liquid, or the mp. if solid, and reft is the following list of bp.s or mp.s, taking into account the part the following list of bp.s or mp.s; taking into account the part to the following list of D.S. on the following the mp.s of mp.s is slentist with, or near to, that of O.S., confirm the identity of O.S. by preparing and determining the mp. of one of the derivatives the indicated. The method of preparation of acetates is described for large S2, of the indicated indicated. The method of preparation of acetates is described for mercesol. page 52, and of methyl ethers on page 182. Brome derivatives should be prepared in the manner described for mercesol. page 53.

Liquids. (See also solids of low m.p.)

B.p.

OH

175\* o-Chlorophenol.

OH

FeCl.: violet (water)

Br Bromo derivative, mp. \$3°.

Bromo derivative, mp. \$3°.

FeCl.: violet (water).

Solids. Mp.

OH

OH

Denzoate, mp 88\*.

FeCt,: violet (water)

OH

\_



HALOGEN-HYDROCARBONS, ETHERS AND ESTERS 141

Вp. CH,CI-CH-CH. 117º Epichlorhydrin.

Glycerol diphenyl ether, m.p. 81°.

Methyl chloroscetate. CH.Cl-COO Me.)

144° Ethyl chloroscetate. CH,Cl COO Et.]

(a) Identify the alkyl radical in the manner de-

scribed under "Esters of carboxylic scids" (page 100). Use half quantities for the hydrolysis, which will be complete in 10 min.

(b) Identify the acid constituent by the preparation of chloroacetamide, mp. 119°, as follows:-To 2 a.c. of O.S. in a t.t add 4 c.c. cone, NH,OH, cork the tube and shake vigorously with cooling until crystals separate. (The period of shaking required will be 1-1 min, for the mothyl ester and 2-3 min. for the ethyl ester.) Filter off the solid, carefully wash it with cold water, dry, and determine the m p.

179" Benzyl chloride. C.H. CH.Cl. Very irritating odour. \$-Naphthyl ether, m.p. 99°.

206° Benzal chloride (Benzylidene chloride). C.H. CHCI. Very irritating odour.

Hydrolysis to benzaldehyde.

Into a 100-c.e wide-mouthed flask pour \$ c.c. of O.S. and 10 c.c. aq. Na.CO, and add some porous not. Fit the flask with a reflux condenser, heat the contents to boiling and continue boiling for 15 min. Remove the flask and cool, -bitter-almond odour of benzaldehyde.

213° Benzotrichloride. C.H. CCl. Very irritating odour. Hydrolysis to bensore acid, m p. 121°.

Proceed as indicated under "Benzal chloride."

Filter, and acidify the filtrate with cone. HCL Cool, filter off the solid and wash it with cold

Crystallise the acid from water, dry, and determine the mp.

Ł,

Bromo compounds.

### Liquids.

B.p.

38° Etbyl bromide, CH, CH, Br. β.Naphthyl ether, m p. 37". (Picrate of this ether, m.p. 104°.)

. m t

MANAGEMENT PT Entroop on A-plan an \$ 111 perce to new-to the title the base fredeside is tended and Visioniament month out fluored to surper surperson Forestetime to be being emission tone families out and 3 ILC. of solid NaDill. Heat to built-diswidne to quite one the next, men & all wheath a drope of did 11cl and 3 H C. of and die (b) To 2 co. alterior in a s s. ach one drop to C (d)

will be obtained. (a) Apply Test (b) under "Chloroform, what lates or no obnoxions odour of phenyl lates and

77 Carbon tetrachlorida, UCI.

order to desirvy the laceyandle. sected, cool and add excess of cone, lift as (carby lamins), Immediately the odour is an duratoni Itaniq lo mobo moixondo gnoris-

too of tol Eurood entitues have NaOH. Heat to boiling in a fume cupband Land to O.H c bas SO to quan and and and MAN

to goth one this 4.1 a ni lodoola .o.o 2 oT (6) bulade no lon toloiv gamesqqa, nice bor-Bunnog or

a c.c. aq. NaOl and one done HOay, pa c.c. u Ma Lt. place 3 R.G. of resortinol and (a) 61° Chloroform, CHCla-

B.p. redniger

Chloro compounds.

hydrocarbons, etc.," (page 144). \*\* Additional in the following lists, see "Additional angular and pale and angular and angular and angular ang to be the leading of or bound four at S.O. H. 1819 og of the undicated. The methods of preparation of derivative angular and entarinab ent to dam ent gammersteb bus gamequiq yo b ang then edi ii oi Saividge yd S.O lo Viiinshi edi mritaco 10 these b.p.s or m.p.s is identical with, or near to, that of OS. whether O.S. contains Cl, Br (page 141), or I (page 142), 11cd. so Il res, noitoesdus estaraqoraque edt ni e.q.m 10 e.q.d 10 sail Determine the b p. if liquid, or the m p. if solid, and refer to Procedure for the identification of O.S.

CHVIN' HYFOGEN ELHERS' VAD HYFOGEN IZLES TOIS THE KI KINDOORH HILM ENGREN IN THE SUCH ALIPHATIC HALOGRY . HYDROCARBONS, ARDAIN

#### Solide.

M.D. 119° Iodoform, CHI., Yellow, Characteristic odour. Apply the tests under "Chloroform" (page 140), when similar results will be obtained.

METHODS OF PERPARATION OF THE ETHERS INDICATED UNDER THE HALOGEN COMPOUNDS IN THE FOREGOING LISTS

(See sections on "Crystallisation" and "Drving of substances."

Pages 16-21.) β-Naphthyl ethers.

In a 100-c.c. wide-mouthed flask place 2 g. 8-naphthol, 1 g. KOH, 1 c.c. of O.S., 10 c.c. of alcohol and some porous pot. Fit the flack with a reflux condenser, heat the contents to boiling and continue boiling for a suitable period of time (10 min. for ethyl, s-propyl, and iso-propyl bromides: 5 min, for methyl, ethyl, \*-propyl, and iso-propyl iodides, also for benzyl chloride and

bromide). Add 20 c.c. water and 1-2 g. solid KOH, cool and shake until solid separates (see note below). Filter off the solid, wash it with cold water, crystallise it from alcohol, dry, and determine the m.p.

Note.

If the b p. indicates that O.S. is an ethyl, n-propyl, or iso-propyl halide, proceed as follows :- Add another 20 c c. water, connect the flack to a water condenser and dutil, changing the receiver when solid appears in the distillate. Continue distilling, adding, if necessary, boding water to the contents of the flack, until no more solid passes over. Allow the water to run out of the condenser, so that the solid, which has collected in the inner tube, will melt and run into the receiver. Filter the distillate, wash the solid with cold water, dry, and determine the m p.

If, instead of a rold, an oil distils over (as will usually occur if O.S. is an iso-propyl balale), extract it from the distillate with ether (see page 21). Dutil off the ether and to the residual liquid add a cold soln, of 1-in, layer of pierie and in a t t, in 3 c.e. alcohol. Stir, filter off the solid picrate, carefully wash it with alcohol, dry, and determine the m p.

Preparation of picrates of \$\beta\$-naphthyl ethers (see note). To tin layer of the other in a t.t. ald I c.e. skyled and beat until solution is complete. Next dissolve fut. layer of peric soil in a Lt. in 3 e.e. hot alcohol. Cool the solutions, then mix and abake. (The perste of Scaphth) | ethyl ether expersion quickly, that of Snaphthyl a propyl ether only after standing for some

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Loughtenit wine, we gut " (Purele of the athet.
                      103. " Empli south (II, CII, CII, I
             23° mar Propyl melula (CII,), CIII.
                                   ( . tot 'd m
  A Stephings other, in p. 37%. (Burese of this other,
                             73. Ethyl uslate Cil, Cil,L.
                                   (.wit du
  A Naphthyl ether, m p. 727. (therete of this ether,
                              43. Hethyl meinie, CH. I.
                                                     ıl b
                                                    'spinby
                                            todo compounds.
 (page 140), when similar results will be obtained
 Apply the tests under "Carbon teltschade.
                       92. Carbon tetrabromide Curt.
                                                    d re
                                                    spiles.
                   6. Naphthyl ether, mp. 99.
     188. Benzyl promide, C.H. CH. Br. Lachrymatory.
               Crum 01 tol lod bus HOA
11, (Used 2 c.c. of O.S. 2 g, policy, 111, 110,
described under " Chloroscette seid " (b), Per
phonoxy acetic acid, m p. 96, in the manner
(6) Identify the acid constituent by prepared
I was of hydrolysis will be complete in 10 mm.
ourd edt tol soitifnaup Mad gniau (001 egaq)
scribed under "Esters of carboxlibe scale
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121. Bromotorm, CHBr. (below).

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Z# I

stranger of in fasiber lysis and the transfer (a) 129. Ethyl bromoscetate, CH Br COO Et. Lachymand Louisido ed lliw estinaen astimis neatw All egay) " miolonold)" nobine sissi edi ylqqA

Proceed as indicated under " Ethyl bromoscetals 144° Methyl bromoscelete. CH.Br COO Me. Lechymidd Glycol diphenyl ether, m.p. 97. 131. Ethylene dibromide. CH. B. CH. Br. (.º67 .q.m

A. Maphthyl ether, m.p. 39. (Picrate of this elbe.

πĴ

HYTOGEN' C' H' [O]

71° n-Propri bromide, CH, CH, CH, Br. Piccate of penaphthyl ether, m.p. 92°. 60. iso Propyl bromide. (CH.), CH.Br.

HALOGEN-HYDROCARBONS, ETHERS AND ESTERS 145

ا مرديءَ المالي

Bp. CH, CH, CH, CHCH.

68° iso-Butyl chloride. (CH<sub>4</sub>)<sub>4</sub>CH-CH<sub>4</sub>Cl.
77° n-Butyl chloride. CH<sub>4</sub>|CH<sub>4</sub>LCl.

100° iso-Amyl chloride. (CH<sub>2</sub>)<sub>4</sub>CH CH<sub>2</sub>-CH<sub>4</sub>CL

106° n-Amyl chloride. CH, [CH,],Cl. Sink in water.

41° Methylene dichloride. CH,Cl.

55° sym-Dichloroethylens. CHCl: CHCl. Unsaturated, but 1 c c will not decolourse

lec of a soln, of Br in CCl, in 15 mm

58\* Ethylidene chloride. CH, CHCl,. 84\* Ethylene dichloride. (sum-Dichloroethane)

CH\_Cl CH\_Cl.

88 Trichloroethylene. CHCl : CCl.

Unsaturated, but 1 c.c. will not decolourse

98° Propylens dichloride. CH, CHCl CH,Cl 120° Trimithylene dichloride. CH,Cl CH,Cl

121 Tetrachloroethylene (Perchloroethylene).

CCl<sub>1</sub>: CCl<sub>2</sub>.

Unsaturated, but 1 cc. will not decolourse

i cc. of a solp of Br in CCl<sub>2</sub> in 15 mm.

147° sym. Tetrachloroethano. (Acetyleno (ctrachlorade) CHCl, CHCl,

158° Glycerol trichlorohydra CH\_CI-CHCI-CH\_CI
159° Pentachlorothane. CHCI\_CCI\_

178° ββ'-Dichloroethyl ether, (CH<sub>4</sub>Cl-CH<sub>4</sub>)<sub>4</sub>O. Solids.

ир.

184° Naphthalene tetrachloride.

Cilci Cilci Cilci

185° Hexachlorosthane, CCI, CCI, Camples-like ofour, (Sublines)

Brome compounds.

Liquids. (Sint in water.)

70° Allyl bromale. Cll,: Cll Cll,Br. Mustard-like odour.
Unsaturated, but I can repaire 20-30 min. for the
decolourisation of j can of a salan of Br in CCl.

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23. tert Butyl chlurale. (C.
                       TR TO CCL* ALTH
                      dood a a a dood
                             to. Vilal opposite.
                               .more
                          40. a-Propyl chloride.
                and Ho
sterest!
         To Fropyl chloride. (CH.), CH-CL.
                                            .d.it
                                       Ploof on water.
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\*spinbit

Chlore compounds.

method described on page 151. work should be consulted. The halogen may be estimated by the

# For further properties and derivatives a more administrated and being a large and the same and t HYDROCARBONS, ETC. NADOLNIO DITABILIA LAKOLTICAN

-Z & phenol, 14 g. KOH. Holl for \$ fir. celycol diphenyl cher from ethylens dibromuse. by drop, until pptn. just commences. Stir, filter, otc. Boil for 5 min,; and 40 e.e. water. To experience heat when the tenfficient alcohol to effect solution, cool and add water, for the respectively, until more, inc.

(Hoth dichlorbydrine are first converted by the KOH w WHO 8 1 hound 3 8 12 minbinding to a lorsoyl 8 (9) HAN 4 1 heard 18 (19 HAN 4)

(a) epichlorbydrin-24 g. phenol, 14 g. KOH. Olycerol diplicing ther from statistic to the anticontrain instructions for or sealisting

ander the name of the ether to be prepared. Boal for the boasts 10 c.c. alcohol and the quantities of phenol and KOH gree bird Proceed as under " f. Naphthyl ethers," using I ca. of 05, Phonyl ethers.

and elimins to all from alcohol, dry, and determine the appropriate the appropriate the appropriate the state of the state solutions just to boiling and mix them. Cool, filter of the man hot alcohol to prevent it from separating out on cooling. Hather pleto solution. Next dissolve 0.3 g. 0f picro seid in just size Meat 0-2 g. of the cther with just sufficient aloniol to effect one

cold sleebel, the following method should be employed Waing to the slight soliding to thinding the sat of meibyl MION.

.q.m eds enimitatels lena.

time.) Filter off the picrate, cardilly wash it with aloohal di-HYPOGEN C' IL [0] \*\*\*

B.p.

CH.

159° o-Chlorotoluene.

Oxidation→o-chlorobenzoic scid, m p. 140°. CH,

162° m-Chlorotoluene.

Oxidation-m-chlorobenzoic scid, m.p. 153°.

Сн.

p.Chiorotomene.

Oxidation-p-chlorobenzois acid, mp 236°.

cı

179° o-Dichlorobenzena. Cl
Solid derivatives difficult to prepare.

...

 $\mathcal{X}$ 

Picrate, m.p. 137°.

Soha. M

M.p., C.
53° p-Dichlorobenzene.

2:5-Dichloronitrobenzene, m.p. 54°. In order to prove that a different substance has been obtained carry out a mixed m.p. determination (see page 14), or apply Test SB (page 171) when a positive test for a native group will be obtained. P-Nino denvariae, mp. ! 133. Oploudensens, CHICL

> Tinpi1 ploro-bydrocarbons. to bette 343.

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E. by preparing and determining the m p. of the dedivative the difficated. The methods of preparation of dedivatives are given Withold with, or near to, that of O.S., confirm the Monthly whether O.S. contains Cl, Mr, or L. If one of these by a or m Subroom, moitosedus estriquique edt ni a.q.m vo a.q.d lo sall Determine the b.p. it liquid, or the m p. if solid, and raid to Procedure for the identification of O.S. :-

### VROMVIJIC HVTOCEN-HADBOCVBROKA

181. Methylene di-iodide, CHala. 12t. u vml ioqiqe CH'l'r 147° 100-Amyl iodide. (CH.),CH-CH,CH,L. 130 u-Butyl jodide, CH.-{CH.]L. IND. 180-1802N 10Q1Q9" (CH1) CH CH T , Ho CH\*-CH 118. sec-Butyl iodide. 101. Allyl iodide. CH.: CH-CH,L. Leck-like odour. 100° ten-Butyl iodide. (CH.),CL. (odo compounds. (Sink in unter.) 219° Glycerol tribromohydrin. CH,Br-CHBr-CH,Br CHB CHB. decomp. 200" sym-Tetrabromoethane (Acetylene tetramom 165. Trimethylene dibromide. CH, Br-CH, CH, br. 142. Propylene dibromide, CH. CHBr CH, Br. 128" n-Amyl bromide, CH. CH. J. Br. 120. iso-Amyl bromide. (CHa), CH CH, CH, Br. 100. m-Butyl bromide, CH.-[CH.], Br. 97 Methylene dibromide. CHalifa iso-Butyl bromide. (CH.), CH-CH, Br. -7.0 CH\*-CH\* 90° sec Butyl bromide. 73. tert Butyl bromide, (CH.),CBr. da

HALOGEN, C, H, [0]

9)]

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Iodo-hydrocarbons.

Liquid. B.v.

188° Iodobenzene, C.H.I.

188\* lodobenzeno. C,11,1.
p.Nitro derivative, in p. 171\*. (No heating is required in the ultration process described on page 140, Allow the nuxture to stand for 5 min. Crystallise from actone and wash the crystals with alcohol)

p.Bromo derivative, m p. 91°.

METHODS OF PREFARATION OF THE DERIVATIVES INDICATED UNDER THE HALOGEN-RYDROCARBONS IN THE FORECOING LISTS

(See sections on "Crystallisation" and "Drying of substances," pages 16-21.)

Nitro-compounds.
To I c.c. or I g of O.S in a dry t.t. add a mixture of 11 c.c. each

of cone. HNO, and cone H.SO. Stand the tt in gently beiling water for 5 min, shaking periodically. Cool, add 10 cc water and shake. Filter, wash the solid with cold water, crystallise it from alcohol, dry, and determine the mp.

p-Dibromobenzene and p-bromoiodobenzene.

In a 100-c, that place 1 c, of 0 S, an amount of iodine about equal us half, to a lad pea and 1 c, becomes. Allow to stand for 10 min, then stard the that, in boiling water for a further 10 min Add 20 cc bouling on Noilly and slade until the liquids become colouries. Cod and shake, then filter. Wash the sold with od-1 water, expanding at from the colouries.

Picrates of a-chloro and a-bromonaphthalene.

To Jam. layer of pierre acid in a Lt, add 3 c.c. of O.S. and shake, until adultion in complete. Cool, add J c.c. of O.S. and shake. Filter, crystallise the sold from alcohol, dry, and the termine the my. Oxidation of chloro and bromotoliuenes to chloro and bromotoliuenes.

benzole acids.

In a 100-cc, with mouthed flask place I e.e. of O.S., 2] g, and KMOo, 40 cc, anter, 3 drups of any Moll and once persus part. If the flask with a redia condense, best the excitout to being and continue being fairly regally for 2 hr. Cc, 4 and pass 50, until any jurgle colour and brown 17st. have disappeared. Filter of the white 17st. of baloyen and, with it will find what pail crystalines from water (or from shooled if the and is notly sprangly table in balony water). By the orytain and determine the my.

2:5-Dibromonitrobenzene, m p. 847. .enoznodomordid-q .ed כאינו בלושלב שמחולי Oxidation-\*- intomolomoto acid, m.p. 251. 28° p-Bromotolueno. B.p. 185°. .q.ıv PiloS Picrate, m.p. 134". 250° «-Bromonaphthalone. Oxidation-re-bromobonzoic acid, m p. 251°. M.p. 28°. 182. p-Bromotoluene. .ccl .q.m ,bion oioxnedomond-m ← noitalaxO 183. w-Iromorojneno CH "ttl q m ,bise nionobenzoic acid, m p. 147". .ensulutomosti-e "[#] "66 q m ,ornternish committe, Callendauray amanda small ned brute est ormaxim mits wedit. Cit many so baltrach erround neatestan soil an benungp Natur derivative, in p. 126°. (No besting b re

HYPOCER C II [6]

135' Remolessen Callalle.

Pint-11 114

411

Brama-kydrocarbons.

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# ALKYL HALIDES AND CHLOROFORMATES

B.p. 151 (b) Preparation of chloroacetamide, m.p. 119°. To 2 c.c. cone, NH,OH add 1 c.c. of O.S.; cool and filter. Crystallise the solid from a small

quantity of water, dry, and determine the m p. ·197° Benzoyl chloride. C.H. COCI. Lachrymatory.

(a) Hydrolysis to benzone acid, m.p. 121°.

Into a 100-c.c. wide-mouthed flask pour 1 c.c. of O S. and 10 c.c. aq. NaOH. Fit the flask with a reflux condenser, heat the contents to boiling and continue boiling for 5 min. Cool, acidify with cone. HCl; cool and filter. Wash the solid with cold water, crystallise it from water, dry, and determine the m p.

(b) Prepare benzanilide, mp 163°, in the manner described under "Benzoyl derivatives" (a), page 217, using 1 c c. of aniline,

# ESTIMATION OF HALOGEN

dethod of Stepanow, modified by Bacon.

Weigh out 0.2-0.25 g. (= w) of the halogen compound and stroduce it into a flask of 300-400 c.c capacity. Add a suitable olume of 98% alcohol ( $w \times 156$  c.c. if Cl is present,  $w \times 68$  c.c. Br, and to X 44 cc. if iodine is present). Fit the flask with a flux water condenser, hoat the contents to boiling, and add adually through the condenser tube, in pieces about the size of pea, a suitable quantity of clean sodium (w x 19-5 g. if Cl is esent,  $w \times 85$  g. if Br, and  $w \times 55$  g if I is present. The idition should extend over at least 30 min ). When all the sodium a been added boal the mixture for 1 hr. Allow to cool somewhat, en add through the condenser tube a volume of distilled water ice that of the sloohol taken. Remove the alcohol by distilling er about } of the volume of liquid. Cool the residue in the flask,

idify with dil. HNO, and add a known volume of  $\frac{N}{10}$  AgNO. lution. Filter off the silver halide, wash the ppt. with distilled ster, and allow the washings to run into the first filtrate.

Titrate the excess of AgNO, in the filtrate with  $\frac{N}{10}$  ammonium ocyanate, using iron alum solution as an indicator (Volhard's

). From the vertice of  $\frac{N}{10}$  AgNO, used up calculate the of hale— at the organic compound.

Ly, . . . . of Ag halide may be dried and weighed.

ost

.boniatdo ed litw saluers railmis meiw ,(461 egaq) (a) Proceed as indicated under "Chloroscotic seid " 103. Chlorosoetyl chloride, CII, Cl COCI, Pungent clour. nim o ni eseigmoo

ed line siavioustitica. Hydrolysis and be " aleriothyd to Lottom to shated" tehm

Dedricesh romann edt ni lasiber lysla edt vlitasbl 94. Ethyl chloroformate. Cl COOEs. Lachrymatory. (69 egaq) " ebinbydna oitsoa " asbuu bodineesb

(5) Prepare propionentitide, m.p. 105, in the mannet propionate in the soln.

a to connersy eds unitimo os , ga ogset ,(a) see'l' Viqa celf. Loniatido ed lliw tiuson nalimis a nedw (a) Proceed as indicated under " Acetyl chloride" (a) 80. Propionyl chloride, CH. CH. COCl. Pungent odour.

Proceed as indicated under "Ethyl chlorolomate," aionyl chloroloma (Proceed as indicated under "Ethyl chloroloma") (Proceed as indicated under "Ethyl chloroloma (Proceed as indicated under "Ethyl chloroloma") (Proceed as indicated under "Ethyl chloroloma (Procedular "Ethyl chloroloma") (Proceed as indicated under "Ethyl chloroloma (Procedular "Ethyl chloroloma") (Procedular "Ethyl chloroloma (Procedular "Ethyl chloroloma") (Procedular " 71. Methyl chloroformate, Cl-COOMe, Lachrymatory, dry, and determine the m.p.

, mixt areq I has lodools strad 2 to crutzing a wash the solid with cold water, crystallise it from then add 10 c.c. rater, cold and all many is the many and a fall many and a fa bas lodidqan-8 g I coalq rasked llama a at

(b) Freparation of franchist acctate, m.p. 70". the liquid, due to the formation of MI, accided to digab edi diguondi bowers) moleo ben-eniw-

an equal volume of aq. FeCla-Pour 2 c.c. of the soln into a t.t., cool, and add 4-5 c.c. by the addition, when necessary, of wake, no longer turned blue, keeping the volume at is i mos equ in poersimmi a furrenom as led alkaline, then boil until a piece of red hims the soln into a dish, add dal NH,0HK and mor 20 lo sqon & libe Al a m rotaw and d oT (u) Acetyl chlorate, CH, COCL Pungent odour,

the identity of O.S. by following the given procedure. erse of three b p. s is identical with, or near to, that of O.S., confirm Il Aqd to tall gain old of the following but and add south raid! -: EO to minulated by tel meleneral

(CHIOROCYBEOXYLES) ACID HALIDES AND ALKIL CHLOROFORIES

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### HELCE III

Althor or Stepane and Transact

A TORRE & SCHOOL NAME & E SCA, DE SERVE "EXPERIMENT

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that, there is promise make the strength of the  $\Sigma$ 

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this of the end one very er. Felly It is its extension of improve extending the following sector of the life of the sector of the life of

that for migraces great on page 1th. If the like often of only increasing a creamed in the a mag determination; if no such proved as induced under "Excess of highest scale" (page 150).

To a 11. - 13 2 e.e. dl. H<sub>2</sub>SO<sub>6</sub> [1 e.e. come you page 154] and heat to d obrar of SO<sub>6</sub> or not, le compounds of a heat plan (reary 184)

"Oxidation of chloro and bromotoluence" (page 149). Oxidation of halogen-acetophenones, Proceed as under eldehyde " (page 62). Oxidation of chlorobenzaidebydes. Proceed as under "Anis-2 : 4-Dintrophenylhydrazones, See page 66. 'eassportee'

> Oxidation - benzole soid, m.p. 121". acetophenone .).

Phenacyl benzoate, m p. 118° (see under "o-Bon" C.H. CO CH, Cl. Very irritating odour. 29. co-Chloracetophenone (Phenacyl chloraus).

Oxidation - p.bromobenzoio acid, m.p. 251. 2: 4 Dinitrophenylhydrazone, m.p. 235.

21. p-Bromacetophenone, Br-C,H, CO CH. Oxidation - benzole acid, m.p. 121.

dry, and determine the m.p. dewith cold water, crystallise from sloobel, donact for \$ hr. Cool, filter off the solid, used

and 5 c.c. of alcohol. Boil under a refux onof O S. J. g. of sodium benzoale, 21 c.c. of sales in a 100 c.c. wide-monthed flack place it I'repairation of phenocit bencome, in p. 115. Call CO-CH Br. Very irritating adout

20. es Bromace tophenone (Phenacel bromuch.)

Og q m bise moznadovoliti-q -- nottelat. 202 d'm anoxempling despone, m p. 265

bet blombe nashlebyde CIC.H. CHO.

PHOS

4 10

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300 gan beine anistrofereiffa q a - enntalur () . 6. 1) Anterphenythy descone, as p. 239. 273. b Lypunn croppenine CI C'H' CO CHE

"El que bisa biogradoredasas a mutchasti 5 1 Bentemphenyldydracon, m p 255

111. or t principantespieplic CICII'CHO

'MI gm Jiga menn aloroldina a nentaligit The g a connentgatengenannel & g 214, \* 1 presignation of Cit. CHO.

ન ગ vyor.tag

-tmotal-negolad tinn politicida-negolad tancitoldef BTOCKE C B' [0] 4:1

#### SCHEME III

#### Compounds containing Sulphur (or Sulphur and Chlorine)

If sulphur only is present proceed as indicated below; if, in addition, chlorine is present, see instead "Sulphochlorides" (page 163).

# O.S. Hquid.

Proceed as indicated under "Esters of sulphuric acid, etc." (page 160);

#### O.S. solld.

If (a) no motal is present, follow the procedure under A.
(b) a metal is present, proceed as indicated under B.

### A. No metal present.

To j.in. layer of O.S. in a t.t. add 5 c.c. water, close the mouth of the tube with the thumb and shake for 15 sec. If any O.S. has remained undissolved, filter. Test the soln. or filtrate with blue litmus paper; if strongly acid proceed as under (1), otherwise as under (2).

- (1) To 2 c.s. of the sq. soln. of O.S. add one drop sq. FeCl<sub>s</sub>. If a violet or red colour is obtained see "Sulphonic social of phonolic compounds" (page 157), otherwise see "Sulphonic social of hydrocarbour" (page 159).
- (2) Apply the general feet for sulphones given on page 163. If the disagreeable garlic-like adout of ethyl mercaptan is obtained identify the sulphone by a m.p. determination; if no such about is produced proceed as indicated under "Exters of sulphone acid and sulphone acids" (page 100).

#### B. Metal present.

- (1) To i.m. layer of O.S. in a t.t. add 2 c.c. dil. H. SO. (1 c.c. conc. H. SO., added to 3 c.c. water. See Note, page 154) and best to boding. Note if there is a pumpent adopt of SO. or pot.
  - If (a) SO, is errored see "Bushkite compounds of ablehydes and hetoma, and sulphorplates" (page 184).
    (b) no SO, is evolved apply Treet 2.

TO CO ITE

Alan | 11) thus similabultation to the a is RO test societies the state of the state of the state of the branchish the branch of the state of the Agula to alon a an it () and a namelled mired along an and and analong A. (621 abed) " stalified adl to making od no largel eX " paint touted to chemical. True the thetilists in the maines described mature. Dual, using a tt as the relatives, until 5 6 ca. of the Mannfult .e.s & sindinos all til fab ber ventiling & segue att Sail said fraunc). Joon , min 6 30) rawmings relief a railed with the male has 1.0 and only only 2 g. CO 3, and some sold and done and and and abl Haber and with mouthed thate pour to a o water, abl J as Land Och Land

# SALTS OF ALKYLSULPHURIO ACTES

(15 % of sq.) landraged to bruoquion ) Sanloon no balos a gaiblisiry White emulsion, or only drops fishe, fragrant odour, with bleuighte nunodusos 1

Mudal	) hanogaos		i .	_
Food in the control of the control o	eq) obygold anolio oiteir siidqlueid	haracte characte diw	noisluu Llio	so esidV/ d 1 i w legonla

b tauoquos elidqineid diiw mobo bnomis. rettid Leanquios siliquisid thire rates odour annual plating concern. The results of the rate of (10 ogad) obydoblalvollas

Yellow colour, odour of cinnamon, with bisulphite compound of cinnamon,

Identification of the sikyl radical-

heating the bisuphite compounds of a few common aroundie the derivative (see pages 65-66). Below are the results obtained by the store by preparing and deferences gainst aritrariab by extraction with ciber (see page 21), identify the attach hepf it to modernia vd bilos it enous to elevable bristodii edi B. Warm 2 g. of O.S. with 20 c.c. aq. Na. CO.; cool Spines " cheloheranone.

" bropionaldehyde." " " 46 .99I anoisi Iqqoiq-n Iqdism .. .. .. .EF1 122° suggests that the liquid is a butyraldebyde. .q.M

\_-

W

page 66) and determine its m.p. examplification the distillate prepare a 2.4 dinition of mort

Liquid floating on the surface of the distillate. 8' C' H' O' [a] 921

# SULPHONIC ACIDS OF PHENOLIC COMPOUNDS 157

# SULPHONIC ACIDS OF PHENOLIC COMPOUNDS AND THEIR SALTS

Only sulphosalicylic acid and its salts, and the salts of p-phenolulphonic acid (sulphocarbolates) are here considered These yield violet or violet-red colour on the addition of aq. FeCl, to their

rocalure :-To 1-in. layer of OS. in a t t. add 2 c.c dil HNO, (1 e c conc INO, diluted to 10 c.c. with water) Stand the t.t. in water at 0° for 2 min.

If there is obtained

(a) a colourless, or faint yellow soln with or without a ppt,

see "Sulphosalicylie acid and its salts"

(b) a red-brown soln, with or without a ppt, see "Salts of p-phenolaulphonic acid " (page 157)

Nore:

A ppt. will be due to the formation of a sparingly soluble sulphate, e g. BaSO. COOIL

ulphosalicylic acid (Salicylsulphonic acid) OH and its salts.

(M p. of anhydrous acid 120°. Hygroscopie) Confirm the identity of O.S by hydrolysis to salicyle send in p. o', as follows :---

Into a 100-cc, bealer pour 5 cc, water and ald 5 cc, cone SO, an amount of O.S. equivalent to I in layer in a tt and me porous pot. Invert over the leaker a 3j in glass famuel. cat the contents of the beaker to lash g and continue being ctly for 3 mm. A sublimate of saley he acad will exclest on the and. Place the funnel in the neck of a separating funnel and ub the deposited crystals into the latter with other. Wash the berred min, three times with about 3 c c. water in order to remove SO. Dutil of the other (me page 21), dry the resides and termine its m.p. OH

tine timedquartement at 10 ath or sulphocarbolates).

no os bas

indicates that O.S. is a salt of methylaulphurle seld, Cil. 80, il. sulphurio sold, Et SO, II, and a positive test for formablebyle -tale to thes a si &O tadt solibition to brainfill about of the a si &O tadt solibidity of H. OZ 13. Hine shinted A (Ct egaq) " stalliteth edit lo coaltus edit no biupil ox " tabru bedrash beman off in stallists the Tree thousand at the Array of the A mixture. Distil, using a t.t. as the receiver, until 6-6 c.c. of da-boil under a reflux condenser for 5 min.; cool. Connect the last boll under a relief a roll of 2 g. of O.S. and some potent blod Into a 100-o.c. water mouthed finsk pour 10 c.c. water, add 3 ca. Los for a con long Oz.H. on Identification of the alkyl radical.

# STILS OF ALKYLSULPHURIC ACIDS

[13 and lineary and the lead of preceding a guidely White emulsion, or oily drops (faint, fragrant odour, with histories 61)

punoduos (79 "Say shedsblatscaltend to stidglasid Arra sqoib benzaldchyde (page 61)

b baseques stidqlesid dire mobe bromis-ratid salioylaidchyde (page 64) Clear, Tellow solin, aromatic odour with bisulphile compound of salicyledchards (record)

Yellow colour, odour of cinnamon, with hisulphite compound of

substance nonmon well a lo shanoquon slidquesid edit Saitasid derivative (see pages 65-66). Below are the results obtained by d bounds of and gaining obligation and action to by extraction with ether (see page 21), Identify the thirty of a mail of the same of a mail of the same of a mail of the same of the same

the liberated althorne or ketone if solid by filtration, or il figal Warm 2 g. of O.S. with 20 c.o. aq. Na, 20, 000 s. cool. ., eycloheranone, \*\* \*\* .001 " propionaldehyde. \*\* .. - methyl a-propyl ketone .. .. •• 113,

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abrablartind as biupil out tada elavaque " eq.m sil enimistel lana (to) eneq.

ed smerbydigasdquimib 4:2 a sudyng stalitzib sås enorg

I little doubling on the surface of the distillate. 3, C, II, O, [CI] v:t

Sulphen.	SULPHONIC ACIDS OF HYDROCARBONS 15				
omide M p.	Sulphonic and (or sail) indicated	Phenyl sater	8-Naphthyl	Suipho	
137*	P-Toluenesulphonic acid CH <sub>1</sub> ·C <sub>1</sub> H <sub>4</sub> ·SO <sub>2</sub> H m-X <sub>3</sub> lene-4-sulphonic acid	Мр. 95°	M p.	М.р. 69	
150*		65*		Oil	
153*	Cult. SO.H	75*		M p. 67*	
	C.H. SO.H	35*	100*	Oil	
154.	o-Toluenosulphonio acad CH -C H - SO H	82*	97*	Ort	
212*	Naphthalene-2-sulphonic acid C <sub>18</sub> II, SO <sub>2</sub> II	98*		M.p. 76°	
D	-10-10 00-17				

Preparation of derivatives of sulphonic acids. (See sections on "Crystallisation", and "Drying of Substances",

P4604 16-21.3

First prepare the sulphonic chloride as follows :- In a porcelain dish place 2 g. of O.S. and add 4 g. PCl, Grind the substances intimately together by means of a postle. If the mixture does not become liquid or semi-solid (which will usually occur if O.S is a free sulphonic acid or an alkali salt) heat on a water bath for 10 min. Cool, wash the contents of the dish with water into a separating funnel and extract with other (see page 21).

Divide the ethereal soln, into two equal portions, distil off the ether from each portion in separate flasks and cool the readless. (Cidorales of p-toluenesulphonic acid and of the naphthalene sulphonic acids solidify, whereas those of benzenesulphonic acid. e-tolurnesulphonic acid, and maxilene-4-sulphonic acid remain liquid.) Proceed with these residues as follows :-

(a) Preparation of the sulphonomede.

Into one of the flacks containing the residue of sulphonic chlorade (which should be melted if it has solutified) pour 10 e a. cone NH OH and star (The sulphonemale thus formed is uscally jellow; the colour, however, dusppears during subsequent operations) Add 20 cc. water and bud down to about } of the volume in order to remove excess of ammonia. then just acaddy with dil. HCL

If the sulphonemale

(I) has dissired completely, evaporate down until erystals experate from a cooled portion. Coal Liter. tack the solid with ould water, and recrystallise it from water.

(2) appears to be maniphie in water, filter, wash the adult with water, and organism it from alcelol. . Dry, and determine the my.

their same in break that prospering for the property of the first property of the prope with sets at these bestelling & money library and Load Predaminants &

TOOL SAND SAND

device plants and a strong and by og as and ministering date and along a gentlemg as and by their stands of a shinderedging outer be ug in all witness h she is united time to a constant dought that to a street of (total many) adminimization to a q m to state the the thirt and and and and and allower and the kulturalennes and to que and sommerate fere manyered EU la mululatura se sol sul sentental

## 8.171K RITERIONIC VEIDS OF HYDROCARDONS AND THEIR

Annuquos eneq a

m E() fad) guitanitail (enomioposned-q in tudicating that () in a pungent a promount of the common and a pungent of the common o

(c) To bein, layer of O in a t t. aid 2 o a, dichronale misture It well with cold water, dry, and determine the up Acidity the filtrate with cone. HCl, filter off the ppt, wash voting of water and litter

sairs na dies othic . Dibite and HOak pa oo & tiba .a.s. a til Idios add valenay T. valew bloo dirw lules add dasw bina And Lond Suisade with a stein at the Br persists after shaking add strong Br soln (10 c.c Br, 15 g. KBr, 100 c.c waler) Soldied of farili ables on the bine 1.1 e ni .8.0 12 ryed, m's 10 analaviups out coalq edus guillod a nI

(b) Preparation of trioromophenol, m.p. 93. (bear birning to notiouber taiting

at to bearn, these simering lo size italia na ot out) ruotoo by dept - nim I sol tother guiliod ni 1.1 oft brate bus Willy agond 2 bbA (egnero of egnand live moleo edt) the yellow soll age. Wall the while the solls is all shall be solled and the soll was formed on heating the salt with HNO., filter.) To 2 th Me ill bine orniq tol test garwollot edt vlqqA bine water, -intense yellow colour, due to the formation of para the fight theore bus 3.4 eds evened, (bemret ed lliw however, a metal such as Ca or Ba is present, a ppt. of sulphale Ma, K, or Zn, is present, a clear soin. will be obtained; if te faus latson a II). bovomer need san emah edt doidw 200 o.c. of water which has been heated to boiling and from Stand the t.t. for 5 min, in a 400 o.c. bester contained blood (a) To a in layer of O.S. in a dry t.t. add I ace cone HAVIstandqualonadq-q tol alast viotamifinoo gniwollol edt viqqA

а, с, н, о, [ci] 891 ESTERS OF SULPHURIC AND SULPHONIC ACIDS

dichromate mixture and some porous pot. Connect the flask to the sloping condenser and distil, using a t.t. as the receiver, until about 7 cc. of distillate is obtained. Test the distillate for the presence of formaldchyde or formic acid, or of acetaldchyde in the manner described under A (page 39) in order to ascertain whether O.S. is a methyl or an ethyl ester.

Treatment of the alkaline residue.

If O.S. is a liquid, acidify 2 c.c. of the alkaline residue with dil. HCl and add two or three drops of aq. BaCl. A white ppt. indicates that O.S. is an ester of sulphuric acid. See list of esters of sulphuric acid (page 161).

If no white ppt, is obtained, or if O.S. is a solid, proceed as under B. "Treatment of the alkaline residue" (page 161) in order to identify the sulphonic acid.

B. Repeat the hydrolysis, but using 5 g. of solid KOH and 25 e.e. of alcohol instead of the 25 e.e. of 20% aq. KOH and boiling gently for 15 min. Add 25 c.c. water and distil off 20-25 c.c. of liquid in order to remove most of the alcohol. Four the residue in the flask into a 100-c.c. cylinder, make up to 50 c.c. with water and proceed as indicated under "No alcohol has been detected" (page 104) in order to isolate and identify the phenolic constituent

Treatment of the alkaline residue (after extraction of the phenol

Acidify with cone. HCl, pour the soln, into a dish and evaporate to dryness on a water bath. Identify the sulphonic acid in the residue by preparing a sulphonamide (see page 159) and determining its m p. (The m p. of p-tolucnesulphonamide is 137°.)

If the whole of the residue be employed, 4 g. of PCI, should be added. Owing to the presence of KCl the mixture will not usually become liquid or semi-solid, and it is advisable to heat it on a water bath for 10-15 min. in order to complete the reaction.

Esters of sulphuric acid.

(3 c.e. completely hydrolysed by lealing with 25 c.e. of 20% aq. KOH for 5 min.) Bp

165° Methyl sulphate. (CII.),SO.

CATTION.

Great care should be taken when using dimethyl sulphate, muce not only is the valant highly Indicator but the liqued is absorbed readily through the skin

A, obtained as under the pass, the condensor into the fash. March 10 cc. varier through the condensor into the condensor into the condensor into the condensor in the condensor

vo. vo. 20, and an analysis of extraction of the controlled to controlled to the controlled of the flast again to the conditions to a state of extern control to the control of the contro

Hydrolysis.

The LiO.co.of. flast, having a short wide neck and flat bottom the will a condensor. Moreover, the contract of the condensor and placed in 155 cc. of 20% aq. HOM, some porous of the condensor and placed in 155 cc. of 20%.

SCIDA.

SCIDA ACIDS only the esters in the following procedure for identification only the esters.

Acrossistence in the lists (pages 16-18) are considered.

OINDHAIDS TO GEA GIOA DIRUHAIDE TO SHATEM

with the control of any historical variables is only one of the control of the co

vivices in the solution of the current with petroloms that [as 2] and 2.41 s.c. and 2.5.41 s.c

(f) Preparative at the place of the framework at the media of the place of the plac

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OFI

# SULPHONES

# SULPHONES

General test.

Mix a quantity of O.S., which would about one-third fill the bulb of an ignition tube, with an equal bulk of animal charcoal. Introduce the mixture into the

-disagreeable garlic-like odour of ethyl mercaptan. Mp.

76° Trional. (Methyl sulphonal)

85° Tetronal. (Et),C(SO,Et), 126 Sulphonal (Me) C(SO Et)

# SULPHOCHLORIDES

To 3 c.c. of distilled water in a t.t. add one drop of O.S. if liquid, or R.G. if solid. Heat to boiling and continue boiling gently for I min. Cool, filter if not clear, and add an equal volume of dil. HNO, and I c.c. aq. AgNO,; shake,—white curdy ppt. of AgCl.
Refer to the last of sulphochlorides (page 169) and identify O.S. by preparation of the derivatives there indicated.

#### SULPHONES

General test.

Mix a quantity of O.S., which would about one-third fill the bulb of an ignition tube, with an equal bulk of animal charcoal. Introduce the mixture into the tube and heat,

-disagreeable garlic-like odour of ethyl mercaptan.

M.p. Me 76° Trional. (Methyl sulphonal)

85° Tetronal. (Et),C(SO,Et),. 126° Sulphonal. (Me),C(SO,Et),.

#### SULPHOCHLORIDES

To 3 a.c. of distilled water in a t. add one drop of O.S. if liquid, or R.O. if rolid. Heat to boiling and continue boiling gently for 1 min. Cool, filter if not clear, and add an equal volume of dil. Rolid, and 1 cc. aq. AgNO; shake,—white cutyly ppt of AgCl. Refer to the hist of sulphochioride (tege 130) and identify O.S. by preparation of the derivatives there indicated.



#### SCHEME IV

Compounds containing N, or N and halogen; also solid compounds containing N with sulphate or phosphate.

If O.S. is a solid proceed as indicated under "O.S. solid" (page 166); if a liquid follow the procedure below.

O.S. Ilquid.

To I ca of O.S. in a t.t. add 5 ca. water, close the month of the tube and invert twice. If OR has

A. desolved completely in the water, test the sold with red himus paper.

If the colour of the paper is

(a) changed to a definite blue, see "Aliphatic and

beterocyclic amines" (page 213).

- (b) unaffected, or little affected, proceed as indicated under " O.S. hould " frace 1771. If O.S. is found not to be formamile, proceed as indicated in Tost 9B (page 171), ignoring (e) under " Tosts with the 4N soln."
- U. not desolved completely in the water, to I a.c. of O.S. in a se add 5 cc dd HOL
  - If (a) a rys. is obtained (misosting that O.S. is an amine, the hydrocklorate of which is sparingly sciolish, add 10 ce. water, best to looking, and, if the mint has not completely desired, continue being with shaking for } man Cook and if any mind is prompt filter. With the sale or fitrate proceed as indicaled to Test & (race 165) continue the lines lost
    - (1) no rea is observed, then the mouth of the 12 and meret team.
      - 11 O.S. 140 (1) described excipitaly as the east, to the

ada apply Test & years Iful conduct the high look If is the lest sector a redbrown colour see sa repulsion is in trulaced. all smite as transfer a orb tre on the 14

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the deat higgs classicates at annies den be man bet ben a ge. sprawed gree as prosper and great their amount with finished and " one (Inchive tient mend were i. a alte) diseased at strong ad-Littlide by Strades male net pared and? &

If the minimum as evening an insure or training the ferifiting in deal and description and Young Your man on An and the attent

execute, tenders, the colours to yendiment, receive the colour Publing maintain ant II) . Anative of by describing and the transfer suitable and dariff, mint & and harden burner burne bached the first in the state of the control of the first of the duming union an al. & O sadt salandan triviou mword qoub to ber Sod a be snamy daval, lantary ants , mm 8-2 ant is s odt adada (page 216). If the birmue paper is unaffected, or hith affected, souling attachtilg to attact " see sean thater of atachting to the blade, a al R.O seolau (byl eg.eq.) " alles muinomah." one , suid studies a communate at angage aumait and to auction and HOak, page care not to touch the glass. If within 30 sec. of adding the bulled, tellul add to fillion and an requer summit box bonsteion to (begand a blod binlado at anoto on it; (000 egat) action of aq. NaOH on picric acid) is produced see Eastern ? ed) to Louisido sadi of ralimis) ruoloo eguaro to wolley escratai as Il 110aN pa .co f lda 11 and 20 to rayel mi g of (1) colourless, pale yellow, brownish, or violet, apply Test I.

H OS grossos a deep yellow colour (e.g. ille) in the solution is if (v. 25er) v colour see Sociou v confirment of the solution of the solution

(b) no metal, follow the procedure below. (bege 113)

"Inward lateld." rebum betaeibni sa besoord "latem a (a) R O.S. contains P1105 'S'O

"nlos Mt ent diw steof" Tobau (a) Purongi (171 eyaq) 88 tesT ni bolanibni sa bound startin to estitite a ed of son bound and al 20 h ; (112 eyaq) " sotenin bine spirit the tests given under "Allyl nitnice Mediated at anoloo ber on test sidt at H Test 8B (page 171), using 5 drops of 0.2 on the sold of plotold mon bestors and (ii)

socime oilogoonled bus oitadqild." ess belidgen & sailedle of " bimpil tastlusor." N' C' H' [O' HYTOGEN] (c) no intense yellow, orange, red-brown, or red colour is obtained, but alkaline respons are overlowd, then to jain, layer of O.S. in a t.t. add 2 c.a. aq. NaOH, and stand the t.t. in gardly boiling water for f min. Remove the t.t. from the water, and hold, for a minute or so, a narrow strip of moistened red littum paper is the mouth of the tube, taking care not to touch the glass. If the colour of the littum paper is changed to a definite blue see "Amides and imides" (page 177); if unaffected or little affected, apoir Test.

NOTE.—In addition to the evolution of NH<sub>4</sub> the bitter-almond educ of benzaldedy would be obtained with hydrobenzanide, (C,H,CH),N<sub>3</sub>, m.p. 102°; an educ somewhat similar to that of benzaldehyde, however, is obtained when benzamide (see page 179) is build with aq. NsOH.

- (d) none of the results described under (a), (b), and (c) is obtained, apply Test 2, unless O S. is a halide salt, sulphste, or phosphate, in which case apply Test 3.
  (2) To i.i. layer of O.S. in a tt. add 2 c.c. dil. H.SO., heat to
  - To j.in. layer of O.S. in a t.t. add 2 c.c. dil. H<sub>x</sub>SO<sub>4</sub>, heat to boiling, continue boiling for j min. and note the odour of the evolved vapours.
     If these smell of
    - (a) formaldchyde (pungent odour), then if the soln is colour-less see "Hexamine" (page 236), or if yollow-brown in address see "Matheless siles" (page 236).
      - colour see "Methyleneaniline" (page 236), (b) Benzaldehyde (bitter-almond octour) "see Derivatives of
  - benzaldehydo" (page 237). (c) neither formaldehydo nor benzaldehyde, apply Test 3.
- (3) To a measured 1-in, layer of powdered O.S. in a day 1-in, 1.1, and 5 c. a distilled vater and beat to builing; if O.S. has not dissolved completely, continues builing with shaking for pim., [If any oil or sold is present, filler.] That the hot soln or filtrate with blue himms paper; note if definitely said or not. (Ghe will be made of this knowledpe later.). Cool and shake well post any separation of solid. [If a solid dissolver completely in the boiling water and does not sprants out at all on cooling and shaking, it will be described for the purpose of the achenes as "mally soluble in water."

### O.S. readily soluble in water.

If the ag, soln, is

<sup>(</sup>i) definitely acid, apply Tost 4.

<sup>(</sup>ii) not acid, test it with red himus paper. If the colour of the paper is changed to a definitelylus see "Amines"

Bill both providing to providing to and the determine brown mar fil all briefe Bireten, an all fi Marat diment dang A dail demand to \$111, 316 at he water on market Lond to

به چه چه موه دو محمد محمد ومده و او دو دو p. 14 part Kelde mongland on mounton a m & O ST 

Let (3, 44). Miny high and mad mad mad be a mind of the distribution of the distributi efeminately at East, decision articled duming a near East II

white or Julius emiliation, proceed se unitrated in ple or soln, am Eactions & (page 210). (a) and imminister red-brown, green, us stated blue calculation of a contract of state of such as the contract of the contract

If the archition of gas is only alightly greater, first pro-(it | soul) A , & noticed one shald out most sadt mant any

(a) a colourless soin, with very much greater evolution of If with the O.S. soln, this produces shoe bloe sucrite barepard ad to soulow

S c.c. dil. Hill as a blank feat) seld, without ahakling, an orquel of only late of the cold appropriate the sent to an C or ( war as 02 m 44 a m bilos edt to tryet mi-f guivlossib to brasish of tam solve. to side pe %12 visionikorqqe the mouth of the Lt. and inverting; do not state, (An Mando vd alm ; EDH Jub to emuloy laupe as bbe ONAV. pe "It to so E of - swolld as bids sucritin to mios a custoff (3)

soluble in dil. HCl," otherwise as "not reachly soluble in dil. Vilen as smades out to secquing out not bodimen as illiw a Dit id galliod ods at viotolquico servicesib bilos eds II MOLE

A secT vique estatifi to mice sidt oT water, boil again, and cool.) If any solid is present, then as (I lds seem bilos a of the ovnd 4.3 off to stratute out 11) 500 ; sim f vol gaileds thiw gailiod sumitance bas gailiod of ted DH Lin as to bba 3.3 a ni B.O to reyed mid of O.S. sparingly soluble in waler.

affected, apply Test & to the aq. soln. dud to hotollann at raded animit out it (202 egaq) soln, is very much greater than that from the blank see "Aliphatia amino-acid" (page 183); if not appreciably different from that of the blank, apply feet 5 unless O.S. contains halogen, in which case apply Test 7.

(ii) not readily soluble in dil. HCl, apply the test under "Carbazolo" (page 208); if a negative result is obtained, apply Test 5 unless O.S. contains balogen,

in which case apply Tost 8.

(5) If no solid is present in the contents of the t.t. reserved in Tost 1 just acidify with cone. HCl, then cool and shake; if any solid is present thick with an equal volume of water and filter before acidifying.

If (a) no ppt, is produced, apply Test 6.

(b) a ppt. is obtained, add R.G. of O.S. to 2 c.c. aq. Na<sub>2</sub>CO<sub>2</sub>, heat to builing, then cool. Add 1 c.c. aq. AgNO<sub>2</sub> and shake.

A dark grey or black ppt, indicates that O.B. is urio

acid; confirm by applying Tost 6.

If no dark grey or black ppt, is obtained, add R.O. of O.S. to 5 c.c. water and heat to boding. Cod, and if any sold its present, fifter. Divide the soln, or fit trate into two approximately equal parts; reserve one of these in case it is respured for the test given under (u)

To the other part of the sola will one drop of Daughe sola.

If (i) a white ppt, is obtained, determine the m p. of O.S.

Mp. 173°-177° milication that O.S. in phenolecte-

toos (lumnal),

189\*-192\* indicates that O.S. is barbices

(u) no fee is preduced, proved as directed in Test & and, if preparity, as in Yest 7, then if this shan the band to the absolutional of

N' C' H' [O' HVTOCEN]

a violet solut a II alos ell lo noitioq by said on eDeT .ps to goth end hhs .2.0

(page 183); otherwise apply Test & shirs oilonedq to sevitavinod " ees bemaido

If there is obtained an orange residue, becoming redained as sources or and rates a no assert bot estrongere and t. C.O to . D. M. nitte (6) Pour 5 co. of a saturated aq. solm of Br into a porcelain dish.

(7) To 2 oc. dil. HCl add B.G. of O.S., heat to boiling with daling I tany other result is obtained apply Teal test, proceed as indicated under "Puring group, (page 343) cooling, and violetred on holding over cone, NH oll (Muratio

Alfa see beniesdo si Agg wolley oleg to estaw a (n) H then cool. Add 2 drops of Mayer's reagent

diphatic and heterocyclic amines " (page 216). to alea on one object in which case one is a a il 20 asslmu 8 Jest Viqqa, benista ei Jqq on (d)

though in a t.t. dissolved in 5 c.c. aq. MaNII., II an immodi the mixture of shaling of amphibio (i. in layer of deaph Na.VO, (ignore any emulaton or slight ppt. formed), Post pe \$10.000 g bbe solar or filtrate add 2 0.0. of or apistate 20 62 bbA mim 5 tol (20 ce, wake, best drown) and drown in the state of th Jone of 1 m. layer in a t.t. of O.S. Continue gratly boiling. Heat the contents of the flask to bolling, and the equite-(8) A. Into a 100-o.c. flask pour 5 c.c. cone. HCl and 5 cc. wald:

the procedure under (b), page 201. noar to that of two or more accept derivatives, follow derivatives (pages 187, 217). If the mp. of 0.5 is It so lo a q m lo stell out of rolor bine &O lo q on out It acctio acid is detected in the dualitate, determine (000 comi) "swimanom vasming ottamora to savit (a) obtained, proceed as indicated under "Arelyl deriver atte, intense, orange-red or red coloured ppt, or solar

(5 ton bas I sat to. 2). bothng and add I c.c. Feblung's soin, (squal solumes he Wall until the mixture is lust alkalued. Heat to (a) not obtained, to 3 e.e. of the acid solut or filtrate sale shallisab adt ni hotsetsis ed bice otters on bluodd

(u) my reduction occurs, princed se imbached mules (ICE stay) " sommed sading late sames() " see thousand at Apy here so expenses and base stamping and II (i) immediate reduction occurs (i.e. blue voluti

041

Repeat Test A with the addition of 1 g, solid SoCI, to the mixture. The immediate formation of an intense, orangered or rod coloured ppt, or soln, with the sikaline famphthol indicates that O.S. has been reduced by the SoCI, and RCI to an aromatic primary sumo compound (NII, in nucleus).

See Section 7 (page 220).

If no orange-red or red coloured ppt, or soln is obtained, apply Test 9 unless the aq. soln. of O.S. was found in Test 3 to be strongly soid, in which case see "Acyl slightatio amino-acids" (page 186).
In a dry tt, place RG, of O.S. one drop of 24% sq.

NaNO, and 2 c.c. cone. H.SO.: warm gently and shake.

If a deep blue colour is obtained see "Derivatives of diphenylamine" (page 208); if no blue colour is produced

proceed as indicated under B.

In a 100-ce, wide-mouthof flask place 1g, of O.S., 15 cc. of a mixture of equal volumes of water and cone. H<sub>2</sub>SO<sub>4</sub> and some porcus pot. Fit the flask with a reflux condenser, beat the contents to boiling and continue boiling for 15 min., periodenally loosening the condenser champ and giving the flask a rotary movement. If O.S. has not dissolved completely, but for a further 15 min.

Detach the flask from the condenser. (Acetic acid, or phenylacetic acid, if present, may be detected by their odour.)

If in the condenser tube there is

(i) a white sublimate, wash it with about 10 c.o., ether into a small separating funed. Wash the othereal soln three times with about 3 c.o. water in order to remove H<sub>2</sub>SO<sub>2</sub>. Distil off the other pages 21, dry the residue, determine the m.p.

so page 21), dry the residue, determine the m.p. d refer to the list of m p.s of sparingly soluble ids (page 75).

, sublimate, allow the contents of the flack to cool somewhat, then add gradually 20 ca. of boiling water. Connect the flack to a condenser, and dusti until 10 ca. of distillate is obtained.

If only a liquid distillate is obtained, test it for the presence of sortic acid or propionic acid in the manner described on page 63.

Should solid be present in the condensor tube, wash it with water into the receiver; Eiter, wash the solid with cold water, dry, determine the mp. and refer to the lists of sparnigly soluble acute (pages 75-79).

at at adt to line adt honesenni yd natio adt ho atendere fore \$ a.c. of the latter through the near of the found mile a day se. buil, alice learned a chi muil eldisont to solaw dount as elengted and cal, gives no blue where on the arbition of starch halele with. quantities of water (3 a.c.) until the lower arround leyer, efter run the breez layer, and wash the upper suffered layer said someone to anyl Jeamed era erayal bothelbure own latest brase or wolfs All 10 c.c. cther (see page 21), liners the stupper, chale, and tunnet unstoppered, states round, and silver to stand for 3 mun. is in indexing a sense of the first sense of a solice sense in the sense of a sense of sense of the sense of se state ONAN pa "ote to no I late found Entiringes a cital Tour 25 c.c. of the 402 mine obtained in Teas 19 steet bydiely

"(Org aSed) .. sorriti...

taking care not to touch the glass. If MII, is evolved, see and so is a distance of the state of the terminal for benedican to ts alkaline. Test for MH, by smell, and by holding a surp (b) To 3 c.c. of the 4N soln, add solid Na 110 mill the mixture

benzoio, or no soid has been detected, son or reno ment and valo bins yns li to ; (102 agaq) " senimanom ynaming stanons to synthetized formon " Bennoys derivatives of annual mand coloured ppt, or soln, is obtained, then if bonzole seal ber vo bor ognero, esentini, intense, intense, orange rol and of her lead of the state of sion is obtained, and I o.c. of the mixture to allaline darph. "man on li ; (woled) " test animescrift" raban botsoibni It a white or pale yellow emulaton is produced at .nim 2 rol basta

of wolls. Suitrorni bas 3.3 eft to strom eft gaired vd and NaW pa %12 to .a.o. I bbe allos Whe diy to .a.o 01 of (a)

Tests with the 4N soln. acida (pages 75-79).

eldde Vguirsqs lo a.q.m to stail edt ot ueld and a.q.m edt eminn age, is obtained, filter, wash the solid with oold water, dry, deterfilter. Cool the filtrate, and just soldiff with cone, HQL II a through the filter; reheat the filtrate and pour again through the O. S. A. pe sailed to see 8 mod (ii) bins (i) in bestead and demonstrate and the second secon attained. (If much solid is present in the filter, and no acid by If the soln, is not perfectly clear, filter until this condition is (.Nh vistamixorqqa ed won lliw .O. H. Cool, and make up the rolume with water to 70 ca. (the

the birs water someon to the bir state of the state of the off Preniment of the residue in the flask. how water. Dry the residue by revolving the end of the tt. over a small faime. Cool, add R.G. of phenol and 8 drope some  $H_2SO_{\rm s}$  and rotate the tt in order to mix the contents. A blue-green or bibe colour, tomed viole-tred by a drop of water, and thus on adding aq. NaOH until the mixture is alkaline, indicates a nitrous scale of the superior of the superior

If acetic acid has been detected in the distillate (Test 9B, page 171) see "Acetyl derivatives of aromatic secondary amines" (page 217); or if benzoic acid has been detected, see "Benzoyl derivatives

of aromatic secondary amines" (page 218).

Metal present.

(Only compounds containing an alkali metal are here considered.)

Procedure :—
To 1-in, layer of O.S. in a t.t. add 2 c.c. water, heat just to boiling,

then coal. If the colour of the soln, is yellow, orange, or red, proceed as indicated under A, otherwise as indicated under B (page 174).

A. If (a) a yellow or corange-coloured solm was obtained, add 2 drops of it to 5 o. water. To half of this duluted solm, add one drop an FeCl.; a deep blue colour indicates that 0.3. is a ferrorganide, a g. K.Fe(Nh). If a deep blue colour in not obtained, to the other half of the diluted solm, add R.O. of powdered FeSO, and shake; a drop blue colour indicates that 0.8, is a ferricyanide, a g. K.Fe(Nh). If negative results have been obtained proceed as indicated under "Bennateer".

(page 52), but omitting the scetone.

The formation of a solid benoy derivative indicates that O.S. is a nitrophenate. (It may be necessary to repeat the procedure, using aq. Na<sub>1</sub>CO<sub>1</sub> instead of aq. NaOII, as 2:4-dimitrophenyl bemoste is readily hydrolized by aq. NaOII.

M.n. of bearonte.

95° indicates that O.S. is a m-nitrophenate. 132° 2:4-dimitrophenate.

142° p-nitrophenate.
(b) a red soln, was obtained, apply the following tests for a nitroprussile (e.g., Na.Fe(CN), NO); —

(i) Add one drop of the red soln, to 2 e.e. water, then add one drop of actions and 2 drops eq. NaOH, —wine-red colour develops, turned violet-red by actio arch.



 $\beta$ -naphthol (}-in. layer of  $\beta$ -naphthol in a t.t. dissolved

in 5 c.c. aq. NaOH)

If a red coloured ppt. or soln is obtained, prepare about 5 ec of a cold, conn. aq soln. of OS. and acidify with glacial acidic acid. Filter off the pptd. acid, wash it with cold water, and dry. Proceed as indicated under "Aromatic amino- and acetylamino- acids" [rays 199].

If no red coloured ppt. or soln is obtained with the alkaline B-naphthol, proceed as indicated under (e).

(e) Apply Test 8B (page 171).

If a red coloured ppt. or soln, is obtained, prepare about 5 c.e. of a cold, cone. aq. soli. of O.S., and scidify with cone. HCI. Falter off the pptd. soid, wash is with cold water, and dry. Proceed as indicated in Section 7 (page 20).

If no red coloured ppi or soln is obtained with the alk-alonhraphthol, proceed as indicated under "Hipperir beid" (page 189); similar results indicate that O.S. is a hippurate. If negative results are obtained, and O.S. contains K, apply Test (a) under "Phthalimide, etc." (page 180). A positive result indicates that O.S. is K phthalmido or K succinitaide. Apply Test (c) page 181); a positive result indicates that O.S. is K phthalimade. a ten'l kurbige by applying Test a A dark grey or black ppt, bullcates that 0.8, le then cool. Add I oc. as AgNO, and shake, Salad to sand , 20 to .0 H Mas OD, av Pa . 32 to T (4)

Microstand group present, and refer to the section att to entire off morroom of volto id (171 egeq) at 8 ard T treserout if bine ,(751 ogwj) & bine C sheef If N is present, apply to the acid (reganied as () !!

If N is absent, examine the soil as under I N tot test onix itselfa

lies the acid from water or alcohol, dry, and apply the off the pptd. acid and wash it with cold water. Crystal-(a) Acidify the remainder of the aq. soin, with HCL, filter

incoron, edulos vigninga et .E.O ii ,10 ,(a) rabnu as bywert oldnios glibert at E.O. it noth, bontable at Agg a (1) H scid and shake.

Fig. 8 c.c. of the cold aq. solu. or filtrate add dil. HC until just ". eldulos vigariada" estwiorito ", eldulos viibsor" beared od live it galdade bas gailooc no lie in two betstades

ton and has roter amiliod ods an ylestologunco beviossib bilos eds 11) solid is present, filter if any solid is present BOH and dissolved dissolved with the shalling with abaing RO H Bullod of fred has roter o.o. o bids at a m. 2.0 to reyal m' & o'T present " (pege 133), otherwise proceed as follows :-

II O.S. contains halogen proceed as indicated under "Metal Procedure for the identification of the acid:

### AMMONIUM URATE AMMONIUM SALTS OF CARBOXYLIC ACIDS, AND

AMMONIUM SALTS: AMIDES AND IMIDES

# SECLION 1

O.S. Hauld.

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(2) no ppt. is obtained, test O.S. as indicated under "Salts of carbox; lie acids," Procedure I (page 87); if this procodure cives negative results with the CaCl, and FeCl. tests, proceed as indicated under " Amides and imides," O.S. solid, A (below).

### AMIDES AND DUDES

Apply the following tests for "Formamide," H CONH.

(a) To 2 a.c. aq. HgCl, add one drop of O.S., heat to boiling and continue boiling for 10 are ,-white ppt. of lig. Ci.

- (b) Heat 1-in, layer of O.S. in a 1.1 .- strong odour of NII.
- (c) To 1 in. layer of U.S. in a tt. add 2 a.e. water, shake to mir, and then add 2 c.c. aq. FeCl.,
- -wine-red colour (viewed through the depth of the liquid); brown ppt, on boiling,

Q.S. solld. (1) To 1-in, layer of OS in a t t aid 5 c e dutilled water and boat to boding with shaking. Cool, and if any solid is present, filter. Tost the soln, or filtrate with blue litmus paper, if definitely

acid proceed as indicated in Section 2. A (page 184), if not acal, aprir Tost 2. (2) To Jun. layer of O.S. in a LL add 2 ec. dil HCL If a brisk efformmenne occurs are "Guanaline carbonate"

(page 152), if there is no effermenace proved as initiated A. Amides and imides indicated by the evolution of NII.

on beating with an NaOll. If O.S. connds of

(a) this present crystals, fallow the procedure under "I'm Indumele" and " Arrancele" (page 178).

(6) a white primite, sparmely mintile in lealing water, apply the tests under "Ozamele" (page 1811

Otherwise carry out a mp decrementation. If U.S. has

(a) me maked at 212", apply the trate trake " ["thehapah. phthalmale, and successful " (sage law, 161 metal at - 1-1 - atts - for said 2 - a a a a a a a a a a a

(20 eyeq) " bins simoidord " bus " bios Cool, and apply Tests (s) and (c) under " Acets 10 c.c. by the addition, when necessary, of water, blue, keeping the volume betwoen 5 c.c. and Vireinamour radge sumili bor lo coold a linu bornation on taut si "nice off ni bornamini bus and manager tuming off unineal soft. then bod mattl the soin, is just all aline, then bod atirities and the sola, is just said. Bairties th the flame, and evened evened and o O.S., and continue boiling gently for 5 min. to bolling. Add the equivalent of J.in. layer in a Pour 10 c.c. aq. NaOH into a porcelain dish and hear

83. Voctemide, CH, CONH. 13. Propionanide, CH. CH. CONH.

indicates that O.S. is methyl carbamato. bine simpol to obridablement tot sast ovilinoq a bas solemedus lydio ei E.O tadi establibri (ce east of positive test for accelebition) in the receiver. Follow the procedure under A A series of the Distil Intell and sucred smos.

the distillate, add 10 c.c. dichromate mixture and tillate is obtained. Rinse out the flash, pour in the condenser tube. Distil until 10 cc. of disto cool somewhat, then pour 10 c.c. water through condenser and continue boiling for 5 min. Albw rules a salem Spilind or lumpit and sand along O.S., 20 e.c. of 20% aq KOH and some porous (6) In a 100-c.c. wide-mouthed flack place I g. of Sulfigge yd estematies lyddom lata lyddo gwraiai

CO, having been evolved from O.S. Distinguish of Surao Affice bornet of the sales eight out the through the larges water for I mun. The a rapid efferremence occurs, and allow the gas to NANO, Attach the La to the cork, best until then self 3 oc. each of till field and 21% aq the cash and place in a O S. to the depth of his contained in another Lt. Detach the the faction breery titles days with about 3 ca, baryta water shods to has add and in nutracine and enastra bae odur Lerrich band ban aten a datm A.d a bill (s.)

Mothyl cathemate, MH, COOMs. Ergli estpourse (Carpone) AR COORT .49 d K

Jennich drapide !! 1:1 1] M p.

114° Ethyl oxamate, (Oxamethane.) NH CO COOEt.

(a) Apply Tests (a) and (b) under "Oxamide"

(page 191) when similar results will be obtained.

(b) Identify the ethyl radical in the manner indicated

under "Esters of carboxylo scids" (page 100), but using 2 g. of O.S. and 10 co. dil. H<sub>2</sub>SO, for the hydrolysis. (Considerable frothing occurs when 20% aq. KOH is employed.) Hydrolysis will be complete in 10 min.

CH, CO
125° Succinumide. | NH
CH, CO

(a) Mix together 3 R.C. of O.S. and twice this bulk of zno dust. Introduce the mixture into an ignition tube and heat. Hold a wooden matchstalk, the end of which has been well moistened with cone. HCl. in the solved vapour.

If the end of the matchstelk is turned a deep red { CH : CH } due to the vapour of pyrrole, | >NH}

CH : CH ) succinimide is indicated. Apply Tests (b) and (c).

If the matchstalk is not coloured red, apply Test (a) under "Benzamide." (b) Apply Test (a) under "Phthalamide, etc."

(page 180), when a similar result will be obtained.

(c) To 1-in. layer of O.S. in a t.t. add 5 c.c. absolutered.

alcohol, heat until solution is complete, then cool.

Add an amount of solid KOH about equal in bulk
to a pea, cork the tube and shake, —ppt. of K

CH \*CO

succinimide, | NK,

succinimide, | >NE CH<sub>1</sub>-CO 128° Benzamide, C<sub>2</sub>H<sub>1</sub>-CONH<sub>3</sub>,

(a) Mix 3 R.G. of O.S. with three times this bulk of dry sods-lime. Introduce the mixture into an ignition tube and heat. A bitter-almond odour of benzonitrile (simular

to the odour of nitrobenzene) indicates benzamide.

Apply Test (6).

. If no such odour is obtained, apply Test (a) under "Succinimide,"

(b) Hydrolysis to beazoic acid, m.p. 121°.

In a 100-c.c. wide-mouthed flask place | g. of -

"(a) ablantations?" Succinitation (d) compounds. Apply Test (b). A green fluorescence is obtained with all three

JOINA TITA I am of the alkaline mixture into a t.t. and fill up and the Malladia at emixture is alkaline. Four colour. Cool, add a few drops of water, then add awold ber a at enutrim oft linn and Viton bas M. SO, and warm. (The smide or imide is thus Mydrolysed to the acid.) Add 2 R.O. of resormed (a) Place R.C. of O.S. in a dry t.t., add 2 drops cond-Phthalamide, phthalimide, and succhamide.

"ON to somut borcopper turnings and 2 drops conc. II, SO.; warm, (b) To 1-in. layer of O.S. in a dry t.t. add a few -vigorous effervescence, orange colour,

NaOCI soln., (all at once from another t.t.) 2 c.c. of cone. heat until solution is complete, then cool. Add (a) To lein, layer of O.S. in a t.t. add 2 o.c. melet.

(Aq. solo; neutral since guantidine is a strong bas.)

and Z drops aq. CuSO, (Febling's No. I sola.);

HOan pe a.o. 2 bbe bas & O to O.S. S conelq 3.3 s al

170° Malonamide, CH, CONH, Proceed as under " Benzanide " (b) (page 179). Mydrolysis to phenylactic acid, mp. 76".

167°. Phenylacetamide. C.H. CH. COMH. mbnu atest adt oale bna (481 agaq) 2 tesT viqqA

"HN 132. Uras. O:C<NH.

water, dry, and determine the m.p. Wesh the solid with cold water, crystellies from and filter.

10 minutos. Cool, scidify with cone. HCl, cool. to gailed emitines has gailed of streeting for Fit the thank with a redux condenser, heat the O.S., 10 o.c. of aq. NaOH and some porous pot

N. C. II, [O, HALOGEN]

N.p. 091

ΔIJ

(page 179). A deep red colour given to the matchstalk indicates that OS, is succinamide, CH. CONH.

(Decomposes on heating into suc-CH, CONH.

cinimide and NH...) If the matchstalk is not coloured red, apply Test (c).

(c) Proceed as indicated above under (a) but using phenol instead of resorcinol. Add the aq. NaOH gradually with shaking until the mixture is alkaline. The red colour of phenolphthalein (destroyed by acid) indicates that O.S. is phthala-

mide or phthalimide. Distinguish as follows:--In a boiling tube place the equivalent of 1-in. layer in a t.t. of O.S. and 10 c.c. absolute alcohol. Immerse the end of the tube in water, which has been heated to boiling and allowed to go just off the boil. Shake for I min., still keeping the tube in the water.

If O.S.

(i) appears practically insoluble phthalamide, CONH.

is indicated. (Melts with CONH ..

decomposition, evolving NH, and leaving a residue of phthalimide.)

(ii) dissolves more or less completely /CO

NH, is indicated

(m.p. 231°).

Confirm as follows :- Cool the alcoholic soln. and pour it into a t.t. Add an amount of solid KOH about equal in bulk to a pea, cork the tube and shake,

-ppt. of K phthalimide. Ozamide. (Sublimes in an open tube.) CONH.

CONH.

(a) In a t.t. place 3 R.G. of O.S. and add 5 c.c. aq. NaOIL. Heat the liquid to boiling and continue boiling for & min. Cool somewhat and acidify with glacial acetic acid.

If the soln, is not perfectly clear, filter until this

-immediate white ppt. (Ca oxalate.) Add 2-3 drops aq. CaCla bondition is attained,

acteristic blue tlame. tlame. CO is evolved and burns with a charand turn the mouth of the tube periodically to the rotating the end of the tube over a small mane, cone, HaSO. Gently warm the mixture of squb & bhe 4.4 tyb a ni . 8.0 lo royal mit o'T (d)

-bing colour. som,); shake, NaOM and 2 drops aq. CuSO, (Fehing s No. ' (c) in a t.t. place 3 it.G. of O.S. and add 2 c.c. aq

18. Chloral formamide. (Chloralamide.) orine present.

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-deep red colour, appearing violet red on shaking Rullod of 150d; Lo. 10 to 10.11 (a) To 2 o.o. seq. MaOH add 3 R.Q. of resorded and cci-cu(on) an cuo:

-white ppt. (Hg,Cl,, due to the preduction of a . nim | 1 and spailind sumitano bas colouriese, Aid I o.o. asp. IlgCla heat to bothing then add dat HCl until the soln, just becomes min. Cool, add a drop of phenolphthalein solitto be being guilted outlines bas guilted of tast (a) I'm layer of O.S. in a t.t. add So.o. aved m. 4 of (a)

mides and imides indicated by epecial tests. deniated of live then talians a male EO Jo 144 w 10 Jokes of \$ In propertitio out Prient Apply Teat (a) minior " Chinconce tie as at " (page 134). Chlorocontamida, Cl CH, COMB, ( operang

 $_{4}0\Omega_{1}\Pi_{1}\left\{ _{HK}^{\ast HK}\right\} 0:HK\right\} \text{ , which we work in$ 

Actionality on for finds became a toda.

title apost in adversary much countries in which the design and the second section in military at minghing their mail 3 3 closes at married 1) To | in Layer of O S. in a t t. a Li S o a, water, heat with יאון שיוטר פוצינולוא בישיווושי tothe and by avolution of CCI, one addition of the little 1)

(c) Preparation of quantidine nitrate, m.p. 214°.

To 1 in. layer of 0.8. in a t.t. add 2 c.c. water, best until solution is complete, then cool. Add gradually 2 c.c. conc. HNO<sub>3</sub>; shake and cool. Filter, wash the solid carefully with cold water, dry, and determine the

Derivatives of phenolic acids (Give a violet-red colour with aq. FeCl.)

M.p. CO-NHC<sub>8</sub>H<sub>4</sub>

135° Salicylanılide.

To R.G of O.S. in a dry tt. add 5 drops of cone. H.SO. Heat for 10 sec over a small fame fabout 1 in. high), cool and carefully dilute with water to 5 c.a. Again cool, and add 5 c.o. of 21/4, ac. NaNO, Add 2 c.c. of this mixture to alkaline f-naphthol (4-in. layer of f-naphthol in a tt. dissolved in 5 c.a. of approx. 2N. NaOH),—red soln, due to the formation of an arocolour.

138". Salicylamide, OH

To \(\frac{1}{2}\) in. layer of O.S. in a dry tt add 5 drops of cone. H, SO., Heat for 10 sec. over a small flame (about 1 in. high), cool and carefully dilute with water to 2 o.e.. Add sufficient solid NaUli to render the mixture atladine and hold a strip of moststend red litmus paper in the mouth of the tube, taking care not to touch the glass paper turned definitely bus, due to evolved NH.

residue (biuret, NH, CO NH CO NH, in I e.s. warm eat eviceals Jood (Liourol otamiidus stidy a bna boylove od liiw HN) solibide taut contactus bottom tiently heat 1-in, layer of O.S. in a dry t.t. until the (v) Pintet (vit. Owe MH CO MH" m b 133.

(rest eged) " sexu "Urea"; if strongly neid, proceed as indicated under "galts of If the eq. coln. of O.S. is nontral, apply the tools given under Urea and its salts

"atlea ed bna sorU " Tohun woled betesibni bosson (a.o. to co. being evolved from 0.S.) proceed as primary samines " (pago 210); if, however, is is tumod If the baryta water remains clear, see " Salts of all baryta

Close the mouth of the lower E.t. with the thumb and without allowing any of the liquid to heavy the upper the and stated out to many out to kinotino out gainvilenant and the street in the street in the street street but and street but in the street in the street but i of Wall and all this third blod birs, and a ONAN pa

until solution is complete, then cool. Add 3 cc. of 21, to the transfer of O.S. in a t.t. add 3 cc. of dil HO, best of (2) reparation of acctons semicarbasons, m.p. 157° (st strongly acid.) Confirm the identity of the base by it of semicarbatide MH, MH-CO-MH, (Aq. solit all Marian

pps, with some effervencence, inclicates that Og is a si and the big of the big of the surface of the surface of the big of the big of the surface of the No. 2) in a Li. just to bothing and all R.C. of O. 20 No commior lemps) and a solidat as a f sault (1) Friends to the standardian of S.S. :-In the cold.

Strouble dilw villbest esgertling altrove steuerprad. . f. EGIOV-ONIEV DILVITATIV TAOV (9)

DIE HOL TAD TO ANNO (\*) CONTROUND REGENTAGE VITEOGEN WIL

I KUILINE

ŧ,

COMPOUNDS EVOLVING N WITH HNO: 185

aq. NaOH and add one drop aq. CuSO; (Fehling's No. 1

soln.).

-violet-red colour.

(6) Preparation of urea mitrate, m.p. 163\*. To j-in. layer of O.S. in a t.t. add 2 c.e. water, heat until solution is complete, then cool. Add 2 c.e. cone. HNO, and cool. Filter, wash the solid carefully with odd water, dry, and determine the m.p.

## Salts of urea. If O.S. is

21

- (i) a chloride, sulphate, or phosphate, prepare urea nitrate in the manner described above under (b).
- (ii) not a chloride, sulphate, or phosphate, dissolve R.G. of O.S. in 2 c o. of boiling 50% acctic acid and add 2-3 drops aq. CaCl.

An immediate white ppt, indicates that O.S. is urea oxalate, mp. 171°.

If no ppt is obtained, to 1-in. layer of O.S. in a dry t.t. add a few copper turnings and 2 drops conc. H.SO.: warm.

Red fumes of NO, indicate that O.S. is ures nitrate, m.p. 163\*,

 Compounds evolving nitrogen readily with nitrous acid only on heating.

#### Aliphatic amino-acids

Refer to the appropriate subsection, i.e. according to whether the aq. soln. of O.S. is neutral or acid.

### Ag. schn. of O.S. neutral.

Aminoacetic scid [Glycine. Glycocoll), NH CH COOH, in p.

(a) To jin layer of O.S. ma ett, add 2 cc. water, sum unta schusen is conjecte, then read, Add 2 dreps of phenolphiladen and a and 2 dreps of NADII, then add as equal volume of neutral advanced lacked the shade to what plannelphiladen and has been added, and then <sup>35</sup>/<sub>10</sub> eq. NADII with just pitch—red reloan despitemen, indicating the neutral character of O.S.

(i) Damire j. a. layer of O.S. in a t.t. in 2 c.c. water. All 2 c.c. aq. CuSO, (Felling's No. 1 min.).—like miner, very much deeper than that of the original aq. CuSO...

(c) Proportion of branch detectors (Hoppins scall, mp.

ot amino-acetic acid.

to the source and also to an equal volume us and to the source of (Tehling's Mo. I soin, larp by drop.

If O.S. is hippure soid a despet blue colour will be obtained with the soin, then with the water, due to the presence

the volume about 5 o.e. by the addition, when necessary.

of water. Cool, filter if necessary.

To the soln, and also to an equal volume of water, add

adul reachton ent ni draseroq ed litre (\*121 ledinale, ledu HOLHO, ance bhe "the Mt. edd 10.c.o. Ol o'T ut Viratnemen reque ambil ber le coole a timu lied ledgest, end bearnt regrot on statie "infoe off at leberton (\*\*Pesseon mefer and info of the statie of the leberton of th

Apply Test 98 (page 171).

Apply Test 98 (page 171).

On 9.2, is hippuric acid a sublimate of benzoic acid (m l

tup 187° Equiv, wt. 179-1.
C.H.-CO.NH-CH.-COOH,
C.H.-CO.NH-CH-COOH,
C.H.-CO-NH-CH-COOH,

(b) Acyl aliphatic amino-acids

(6) [bryans the bentoyl derivative, m.p. 180°, in the man the described under "Aminoacetic soil" (5) but is g. of NaHCO.,

"the action of the major (a) and (b) and (b) when the major when t

4. taninomuccinio acid (4. tapartio acid), (CHUM.)-COOH,

#### SECTION 3

#### COMPOUNDS YIELDING DIAZONIUM SALTS WITH DIL. HCL AND AQ. NaNO.

(These compounds will have been indicated by the formation of a red- or brown-violet dye on the addition of their diazonium or tetrazonium salt to alkaline 8-nanhthol.)

Follow the appropriate procedure, i.e. under "0.8. liquid" (below) or under "0.8. solid (page 185). The general methods of preparation of derivatives are given on pages 217-8, and the method of determination of equivalent weights on page 218. The equiv. with green have been calculated from the formulas using the following atomic weights:—C=12 00, H=14008, O=1640, N=1401, C=3546, Br = 7992.

### O.S. tiquid.

Prepare and determine the mp. of an acetyl derivative, then is refer to the appropriate list (according to whether halogard absents or present) of mp. so facetyl derivatives in order to identify the amine. For confirmation of identity, repare the beauty derivative and determine its mp., or determine the equir. wt. of O.S.

### Halogen absent

Acetyl dermatica M p.	Amine indecated NH.	D p. of	Benzoyl deriu, M.p.	Equity, set, of
62.	m-Tolaidine OCH :	199*	125*	107-1
84*	o-Aniaidine OCH	21s*	64*	123-1
101*	Methyl anthramilate, See	page 183.		

A rebnu subcoord ed wolld, follow the procedure ander A. If the aq. soln, of O.S. is definitely acid, proceed as indicated bilos .2.0

(.nuoloo nword-wolloy a tine saria entitine (o. Toluidine gives only and stir for about 10 soo. Old one or two days of lands and stir for about 30 solid with water to 10 co.), If the amine of the amine of the solid soli and then dip the end of the rod into 5 o.c. water contained in a the touch the surface of the amine with the extreme tip of a glass rod, sult : swollot as embiniot-o most badainguists ed yam emlinA 87° o-Chloroaniline .407 .00I

c-121 72" m-Chloroaniine .071 CI bicseut "HN CH, 133. 5-Ymmo-b-xkw 1-151 CH 13.0 1922. Delipeneration .221 ·LEI \*HN "RS 122. Plumber 21s. .:61 121 "HO .951 .691 26 113. at-CI warms Z C H let uvroceal 67.1

A. Au, soln, of O.S. not acid,

Determine the m.p. of O.S. and refer to the appropriate list of m.p.a, i.e. according to whether halogen is absent (page 189) or present (page 195). If one of these m.p.a is identical with, or near to, that of O.S. confirm the identity of O.S. by following the given procedure, or by preparing one of the derivatives indicates

Norras.

(i) When O.S. has a m p. identical with, or near to, that of both a free amine and an acetyl derivative of an amine, either the procedure described under "Acetyl derivatives of aromatic primary monamines" (rage 200) may be carried out, or an attempt to prepare an acetyl derivative may be made (see page 217). In the former case a positive test for accide acid in the dustile will include the document of an early derivative, and the second of the contract of the contract of the derivative, and the contract of the derivative of the contract of the derivative of the derivative

(ii) In the case of a free amine, a determination of the equiv. wt. will provide additional evidence of identity.

Halogen absent. M.c.

M.p. 25° Methyl anthranilate

COOM!

Orange-blossom odour.

Proceed as indicated under "Esters of carboxylic

acide "(page 100) but using half-quantities, when a positive test for methyl alcohol will be obtained. (Hydrolysis will be complete in 10 min.) Acidify the alkalme residue with glacial accide acid, filter off the pptd. anthramilie acid, wash it with

cold water, dry, and confirm its identity by determining its m.p. (144°).

Acetyl derivative, m.p. 101°. Benzoyl derivative, m.p. 100°.

NH,

43° p-Toluidme,

Powerful characteristic edour.

Acetyl derivative, mp. 148°. Benzoyl derivative, mp. 158°. Equiv. wt. 107-1.

m p. 158". Equiv. wt. 107-1. 46" Formanible. C.H. NH-CHO. Faint calour.

(e) To 1-in. layer of O.S. in a dry s.t. ald 5 drops cone. H.SO. Gently warm the mixtures by

characteristic blue flame. the flame, CO is evolved, and burns with Ulabiboing edut edt to diuonr edt mut bna rotating the end of the t.t. ever a small fan

chloride, etc., as described under" Benzoyl deriva mixture is alkaline, then adding 2 o.c. bemol Il Ilon HOan succepts gained at 163°, by adding bilinexned eneque dash eat at biupil laubison eat off benistdo ed lliw atluser evilizoq nedw (6,83 HgCl, and AgNO, tests for formic acid (see pa distillate is obtained. To the distillate spely t condenser for 5 min, then distil until 5 c.c. O.S. and 15 c.c. dil H.SO. Boil under a refi (6) In a 100-c.c. wide-mouthed fleek place I &

Acctyl dorivative, m.p. 159°. Benzoyl denvative 50° c-Naphthylamine. \*HN tives " (page 217).

\*HN m.p. 160°. Equiv. wt. 143-L.

OV-HN m.p. 151. Equis. wt. 123-L. Accetyl derivative, m.p. 127°. Bonzoyl derivative, O-CH\* onibisinA-q "73

entibilities o lytook \*48 DIF-TEN enibiulot-m lytsoA .ca

69. Benzoseine (Ethyl p-amhochenzuste).

\*11N stomatic primary monanthes " (page 200). Increased as indicated under " Acetyl derivatives

13000

### AROMATIC AMINO COMPOUNDS

101

M.p. Proceed as indicated under "Methyl anthranilate" (m.p. 25"). M.p. of p-aminobenzoic scid 186°. Benzoyl derivative, m.p. 148°.

105° Propionanilide. C.H. NH-CO-CH, CH.

Proceed as indicated under "Acetyl derivatives of aromatic primary monamines" (page 200).

The blad distillate apply Tark (a) from 691 for propriories

To the distillate apply Test (c) (page 63) for propionic acid, when a positive result will be obtained.

NH-Ac

110° Acetyl e-toluidine.

3]

Proceed as indicated under "Acetyl derivatives of aromatic primary monamines" (page 200).

112° \$-Naphthylamine. NH.

Acctyl derivative, mp. 132\*. Benzoyl derivative, mp. 162\*. Equip. wt. 143-1.

114° Acetanilide. C.H. NH-CO-CH.

dry, and determine the m.p. NH-An )

(a) Proceed as indicated under "Acetyl derivatives of aromatic primary monamines" (page 200).

(8) Prynamion of phromocentamilide, m.p. 167°. In a 100-ca. flask place 1 g. of O.R. and 6 c.e. glacial acetic acid, heat until solution is complete, then cool. Add 8 c.e. of a solz. of 1 c.e. bromine in 9 c.e. glacial acetic scid, and allow to stand for 15 min. Add 50 c.e. water and abake. Filter, wash the solid with water, cyteallies from alcohol.

127° Acetyl p-anisodne

127° Austyl 4-amino-m-zylene OH-As

Yield red dyes with alkaline p-naphthol in Test & (page 165).

он. си.

Proceed as indicated studer "Acres) destructions of accommand primary monancies " (page 200).



M.p.

3]

NH-An

135° Acetyl p-phenetidine. (Phenacetin.)

(4) Proceed as indicated under " Acetyl derivatives of

aromatic primary monamines" (page 200). (b) Preparation of 3-nitro derivative (yellow), m.p. 103°.

In a 100-c.e. flask place 1 g. of O.S. and add a mixture of 1 c.c. cone. HNO, and 9 c.c. water. Stand the flack for 5 min. in boiling water and shake round periodically. Add 20 cc. water, cool and shake. Filter, wash the solid with

water, crystallise from alcohol, dry, and determine the mp.

139° Acetyl 2-amino-p-xylene. NH-Ac

Proceed as indicated under "Acetyl derivatives of aromatic primary monamines " (page 200).

COOMe

142° Orthocaine (Methyl m-amino-p-hydroxybenzoate)

(a) Dissolve R.G. of O.S. in 5 c.c. water. To the cold soln, add one drop ag. FeCl., -violet-red colour, rapidly changing to brown,

(b) To 1 in, layer of O.S. in a t.t. add 2 c.c. aq. NaOH and shake.

-0.S. dissolves readily, owing to the presence of a phenolic group, Benzoyl dehvature, m.p. 242°. (Dissolve I g. of O.S.

in 5 c.c. cold acctone, add 2 c.c. benzovi chloride and allow to stand for 5 min. Filter, wash the solid with acctone, dry, and determine the m.p.) 147° Phenylures. C.H. NH-OO-NH.

(a) Heat 1-in. layer of O.S. in a t t. above its mp., -strong odour of NH, (Difference from sorty)

AI

N' C' H' [O' HYTOGEN]

To 1-in, layer of O.S. in a t.t. add 2 ce d (C'H'-NH) CO' m.p. 238". (b) Preparation of diphenylurea (artanulue),

and then with water. Crystallize from slooked and filter, Wash the solid, first with dil HO. Allow to cool, add 10 c.c. boiling dil HC, clair litmus paper in the end of the condenser tube. on detected by holding a strip of moistened for but hos bevious in HM , bilos as tuo sestateque continue boiling for 10 min. (The carbanite has tailed ot 1.1 edt to sinstnee edt teeH. and 2-ft, tube (to serve as an air condenst). treshly distilled amiline. Fit the t.t. with a mit

oA-HN dry, and determine the m.p.

148° Acetyl p-toluidine.

(WE eyed) " semimenon vasming oitsmore to sortievinsb lytech." rabau botsoibai sa boscord ence from phenylures, m.p. 147°.) Does not yield NH, on heating above its m.p. (Differ

NH-YO

159º Acetyl a-naphthylamine.

evidential lytooh" about betselbni as becoord (a)

Randan-Altion " tobau betalbai es bescorf (b) Preparation of 4-bromo derivative, m p. 193". (002 eyeq) " seminanom vasming ottamora to

acetic acid for the solution of O.S. bilnel 2 to .0.0 8 gales tud ,201 egeq (6) " enims

ov-HN

162° p-Aminosoctanilide.

(i) To 5 c.s. of the deluted sole , at 1 c.s. of Dinks with water to 100 c.e. heat to boiling and centime boiling for I man. (a) To 1-in. layer of O S. in a t.t. ald 5 e.s. dil IlCh.

معتراه سنهم طف

FeCl, then 1 c.c. of a saturated aq. soln. of H.S.

-violet colour.

(ii) To 5 o.c. aq. NsOH add 1 c.c. of the diluted soln, and 1 c.c. of a saturated aq, soln, of phenol, then add 1 c.c. dill. NsOH soln, (1 c.c. conc. soln. diluted with water to 10 c.c.) Allow to stand for 2 min., —deep blue colour.

(b) To 1-in. layer of O.S. in a t.t. add 2 c.c. dichromate mixture and heat to boiling,

-pungent odour of p-benzoquinone.

(e) Proceed as indicated under "Acetyl derivatives of aromatic primary monamines" (page 200), when a positive test for scetic acid in the distillate

### Halogen present.

### AS HALIDE SAIT.

M.p.

155° Proceine hydrochloride. (Nevocaine. Hydrochloride of diethylaminoethyl-p-aminobenzoate.)

COO-CH, CH, N(Et), HCl.



will be obtained.

(a) Dissolve R.G. of O.S. in 2 c.c. water. To the cold soln, add one drop of Mayer's reagent, —white ppt., dissolved by adding an equal volume of dil. HCl.

(b) Dissolve R.G. of O S. in 2 c.e. water. To the cold soln, add one drop of iodine soln. —red-brown ppt.

-red-brown p

#### NOT AS HALLDE SALT.

Cl present.

M.p. 70° p-Chloroaniline.

Acetyl derivative, m p. 178°. Benzoyl derivative, m.p. 192°. Equiv. wt. 127.5.

of sect ! I have it's cubl soln, silt 1 co, ill litt and then 1 co. (i) Dissolve It Ct. of O.S. in 10 cc. water. To the -: eyen Binampos ags Aq amme ants to Alla is indicated. Confirm the identity ted to brown, a selt of p-phenylenelles. (a) a deep green colour, which rapidly changes through If there is obtaunal solution is complete. To the cold soln, add one drop of aq-To 5 c.c. water add R.C. of O.S. and shake or heat until Procedure: 3. Aq. soln. of O.S. definitely acid. toon again, " sominanom Traming oitamora to Proceed as indicated under " Acetyl derivatives 167° Acetyl p-bromoaniline. OV-HN m.p. 202", Equiv. wt, 172-0. Acetyl derivative, m.p. 167°. Benzoyl derivative ed" p-Bromoaniline Br present. ton eyeq) " sammenom vraming oldemore to ertiarinab lytsoA " rabnu botsoibni aa booorti 178 Acetyl p-chloroaniline OA-HN 87. Acetyl o-chloroaniline DA-AN .q.14 N' C' H' IO' HVI'OGENÌ

961

Δij

-speke test colour.

(ii) Dissolve R.G. of OS. in 5 c.c. water. To the cold soln, add 1 c.c. aq. NaOH. 1 c.c. saturated phenol soln, and 1 c.c. dil. NaOCI soln. (1 c.c. conc. soln, diluted with water to 10 001 -deep blue colour in about one min.

(iii) To 1-in, layer of O.S. in a t.t. add 2 c.c. dichrom-

ate mixture and heat to boiling. -pungent odour of p-benzoquinone.

b) any result other than that described above under (a) follow the procedure given below. (With FeCl. salts of benzidine and tolidine yield a blue, or persistent green colour, and salts of g-naphthylamine. but not of 8-naphthylamine, develop a blue-violet t.ruoloo Procedure :-

To 1-in, layer of O.S. in a t.t. add 2 c.c. ac. NaOH. close the mouth of the t.t., and shake vigorously for 1 min

If (i) O.S. has dissolved completely, add 1 c.c. placial acetic acid, cool well and shake pigoroutly.

If a ppt, is obtained see " Aromatic aminoand acetylamino-acida" (page 199).

(ii) an oil or emulsion is produced, see "Salts of aromatic primary monamines" (page 198). Also see note below

(iii) a solid is present, then if the dispotised soln. of O.S. gave a red dye with alkaline 8-naphthol see "Salts of aromatic primary monamines" (page 198), or if a brown-violet dye was obtained see " Salts of benzidine and tolidine " (page 198).

Also see note below.

#### Norz

If an acid radical has not been already detected add 10 c.c. water to the alkaline mixture, shake and filter. To 2 c.c. of the filtrate in a tt add I ce dil II SO, and 2 R.G. of powdered FoSO.; shake until the latter has dissolved. Pour (from snother Lt.) 2 c.c. of cone. H.SO, carefully down the side of the tube. A dark brown rung at the junction of the bounds indicates a mirate.

بعديده

( but many) " mountains " tone " mountains " mes. The dute these at the sites minute, want is sith will solid, but שיביון ב שיין פורב צים מים ל וויינו פינים בווים וויום שיווי שיווי ( ויים נווים bellind minimit + (130 4 5 544 fere Landed on said Alon da Tree & Liber o. Smare Laute list a.m. water stell lib are to Bull. miliand has and send by the

despects the energy decivation.

whiled mitting and this dies deed tottes at fainte (out many) manufaces are chose ingrituir, the tent thousthed under " Mide "

Since the major to every characters to end in e.g on mile would

file Lt saint seametin france momentum (jeges 1st, 211). It southwittels It land to a q m to (amount to smooth at mangalan m.p. and refer to the appropriate list (according to whiches test it for the absence or presence of halogen. Deletinise the If O 3, contains balogen, receystallise the scotyl derivative and

(VIE egal ow) onima od to evitavirab lyton an

page 31) and duttl off the solvent. (See note below.) Prejace cold water and dry; if a liquid, extract with 10 a.a. ather (are stir. If the liberated amine is a solid, filter is off, wash is with to cool somewhat, then add 20% KOR until alkaline, cool and necessary adding boiling water, until solution is complete. Allow water,) Add 2 g. of O S., continue gently bothng and sin, il (Meny sales of amines are much more soluble in dil. Hel than le Your 10 c.c. dil. HCl into a 100 co. beaker and heat to beature Procedure for the utentification of the amine.

Saits of aromatic primary monamines.

'020 '02022302 malos ne si .8.0 li nistrones ot robro and apply Tost (b) under A (page 87), in addition, when necessary, of water, Coll the volume between 6 a.a. and 10 ac. by the the soln, is just no longer turned blue, kerput in leersman plinstanemon roqeq sumril ber the soin, into a dish and boil until a piece of add dil, NH,OH until just alkaline, Jum and o'O'H Lib dire etartia enitsala edt lo If no ppt, is obtained, socialy the remained A white ppt, indicates an oxalate.

seid, heat to boiling and add 2-3 doys at 2 c.c. of the slkaline filtrate with glacial axes Line honicido at thusan evidence a II

Ш

Aromatic amino- and acetylamino- acids

Determine the m.p. of O.S. and follow the procedure under the ompound whose m.p. is identical with, or near to, that of O.S.

COOH NH. M.p. 144° o-Aminobenzoio acid (Anthranilio acid).

> Preparation of acetyl derivative, m p. 185°. To 1-in, layer of O S. in a t.t. add 3 c.c. acctone, shake until solution is complete, then add 1 c.c. acetic anhydride. Allow to stand 2 min., then add 5 c.c. aq. NaOH in order to neutralise some of the acctic acid formed (the soln, should still be acid: if not, just acidify with acetic acid), cool and shake.\* Filter, wash the solid with cold water, then add it to 15 c.c. boiling water contained in a beaker. Stir, adding if necessary a few drops of glacial acctic acid, until solution is complete. Cool, filter, dry the solid and determine its m p.

COOT 174° m-Aminobenzojo acid.

> Preparation of acetyl derivative, m.p. 250°, To 1-in, layer of O.S. in a t.t. add 5 c.c. water and I c.c. sectio anhydride. Close the month of the t.t. and shake vigorously for 2 min. Cool, then proceed as described under "o Aminobenzoio acid," commencing at the asterisk,

COOII NII. A 185° Acetyl anthranilic acid. 186° p-Aminobenzoio acid.

> Proceed as indicated under "Acetyl derivatives of aromatic primary monamines " (page 200). If (a) acctio acid is detected in the distillate, add to



roulk will be abtained. Wash the solid with cold water, then add it to 50 a.c. boiling water contained in a beaker. Continue boiling with stirring until solution is complete, then cool. Add 10 a.c. aq. NaOH, cool and stir well. Pilter off the pybd. amine, wash it well with cold water, dry, and determine the m.n.

M.p. 50° indicates that O.S. is acetyl α-naphthylamine.

112° ", " acetyl β-naphthylamine, (ii) no solid hydrochioride has separated, continue bolling for 10 min., then distill until 7-8 c. of distillate is obtained. Test the distillate for the presence of acetic acid in the manner described on page 68. If O.S. is an acetyl dirivatire a positive result will

be obtained.

In order to distinguish between two or more acetyl derivatives with m.p.s. close together, or for further confirmation of the identity of O.S., proceed as indicated mades (b).

indicated under (b).

(b) Add to the contents of the Eask aq. NaOH until the mixture is alkaline, cool and take. Prepare a benzoyl derivative of the liberated smine by adding 2 oc. benzoyl chloride, ctc, as desembed on page 217. Determine the nn.p. and refer to the following appropriate last (according to whether halogon is absent or present) of mp. a of benzoyl derivatives.

Benzoyl derivatives of aromatic primary monamines.

#### R-NH-CO C,H,

### Halogen absent.

If the determined mp. is near to two of those given below, obtain (or prepare from the amine as described on page 217) one of these bemoryl derivatives and carry out a mixed m.p. determination (see page 14). Mr.

64° Benzoyl o-anizidine, 125° , m-toluidine, 140° , 2-anino-p-xylene, 143°

143° " o-toluidine. 154° " p-anisidine. 158° — p-anisidine.

158° " p-toluidine. 163° " anilme (Benzanilide).

173° " p-phenetidine.

92° " 4-amino-m-xylene.

A' C' H' [O' HYTOGEN]

THEFT IT

.cci Fenzori e chionendine (a Chlombenzaulide).

470 THE SHIMM 103.

203. Henryl p-bromosniins (p-Bromobenzaniids).

Tost 93 (page 171). socion are those detected by the age dys test after bydrolysis in cluded in Section 3 (page 189). The amilides included in the Amiliales of formic, scetic, propionic, and benzoic scale are m-Anilides

and saired the continue the dentity of O.S. by following the given m.p.s. If one of these m.p.s is identical with, or nest to, the Determine the m.p. of O.S. and refer to the following list of --: sintoxof.

For m.p.s of other anilities see under the carboxylis acid dekeded procedure.

Succinanilida. сн'со-ин-с'н'

alkalme soin, into a t.t. and fill up with water, until the mixture is alkaline. Pour I c.c. of the add a few drops of water, then add a NeOH until the mixture is a red-brown colour, Coolresortinol and 2 drops cone, H.SO., Gently hear To R.C. of O.S. in a dry t.t. add twice the bulk of сн\*со∙ин∙с\*п\*

scetic soid, Meat to boting under a rettax conand 21 c.c. each of social sake place 1 g. of O.S. and gladish Preparation of acciantiste, m.p. 114". 238 Carbandide (Diphenylurea). (Cata-MH), CO.

-yellow-green fluorescence.

.A6 180T m

dn

ar.

flack a rotary movement. Add 30 c.o. boiling water, boil for 1-2 min, then filter. (Sool, filter ally losening the condenser clamp and graing the denser and continue boiling for 15 min, periodio.

AMINOPHO.S. in a dry t.t. add 5 drops M.p. CO.NH.C.H. itly warm the mixture by rotat-245° Oxanilide. tube over a small flame, and

CO-NH-C.H. f the tube periodically to the (a) To 1-in, layer of ved and burns with a charactercong. H.SO., Ger.

ing the end of the nouthed flask place 1 g. solid turn the mouth o c.c. alcohol and the equivalent flame. CO is evolvit.t. of O.S.

istic blue flame. under a reflux condenser and

(b) In a 100-c.c. wide- 5 min. To 2 c.c. of the liquid KOH, 5 c.c. water, , glacial acetic acid until the of 1-in, layer in a idd 2-3 drops aq. CaCl..

Heat to boiling | malate.) continue boiling for from the flask addiols

soln. is acid, then es of p-aminophonol, 2:4--white ppt. (Ca shenol are here considered. Aminophe ated by the deep red or deep

(Only hydrochlorides and sulphs diaminophenol, and p-methylamine, I shake or heat until solution

OH

ŇН.

These compounds will have been indie drop of aq. FeCl, and note brown sola, obtained on shaking the within 15 sec. Procedure for the identification of O h' is obtained, apply the tests

To 5 o.c. water add R G. of O S. ar-is complete. Add to the cold soln, our within 15 sec., proceed as any colour produced immediately, o aminophenol."

If (a) an immediate wine-red color, under (a) and (b) is obtained,

under " Amidol." (6) a deep violet colour develon

indicated under "Salts of 1. NH, (c) neither of the results describes of apply the tests under " Met al.

, c.c. cold water. Add the Amidol (Hydrochloride or sulphate) 2:4-diaminopher of AgCl if O.S. is a hydro-

(a) Dissolve R.G. of O.S. in I limewater and abake. soln, to 2 c.c. aq. AgNO, ring to deep blue. -deep red colour (with ppt. chloride).

(b) Add R.G. of O.S. to 2 c.c. -yellow soln., rapidly chan

shake until solution is complete. Add 2 s.c. acotte To tale, layer of O.S. at a at E.O to rotal all or byenet mp. 240.

(s) Preparation of monoacedyl derivative of p-methylamino--pungent odour of p-benzoquinone.

Builtod of Jead bes emixim

(b) To 1 in. layer of O.S. in a t.t. add 2 c.c. dichromate Donialdo ed lliw enimesoriin a not test eviticog a medw (802 eged) "sonime Trabnocod" Teban moislume ed! Tol bedinesh remam edt ni erutzim edt tavrl .(\*861 q.m) -ppt, of pale-yellow, matted needles of nitrosamine, NaNO, and shake,

ps %12 lo .o.o 2 bbA .obleton is complete. Add 2 o.o. of 21% aq (a) To 1-in. layer of O.S. in a t.t. add 2 c.c. dil, HCl and

"ITO HN Metol (Sulphate of p-methylaminophenol,

·q.m

wash the solid with cold water, dry, and determine the for I min. Cool, add 5 c.c. water and shake. Filter, dratide, heat to boiling and continue boiling with shaking To lim, layer of O.S. in a t.t. add 3 c.c. acetic and 4m horsing on man derivative derivative of p-aminopheno, and

and determine the m.p. off the yellow ppt., wash it well with cold water, dr. soln, add 5 c.c. conc. NaOCI soln, and shake, Filer warm or shake until solution is complete. To the cost DH dh as & bhe at a m a CO to sych mi f oT

(c) Preparation of quanomechloroimide, m.p. 63. -bungant odour of p-benzoquinone. Eniliod os sand bas emittim

(6) To 1-in. layer of O.S. in a Lt. add 2 c.c. dichromas mim I snode ni eqoloveb tuolos enid qosbed byenet com: spepa

crish solm, sold I c.c. sep. MaOH and I c.c. of saturable (a) Dissolve R.G. of O.S. in 20 co. water. To Sea of the

hundonemb q la stief.

AMINOPHENOLS 205

# anhydride and shake vigorously until crystals separate. Filter, wash the solid with water, dry, and determine

3]

the m.p.

Dibenzoyl derivative of p-methylaminophenol, m.p. 175°. (For method of preparation see page 217).

### \* KOLLORS

'ONDN 'OF ANY KETTOIA ENIGTRION MILH DIT' HOL COMBONNES LIETDING Y MHILE OR

and the production of a red ppt. of Cu.O) indicates that Od is Immediate reduction (i.e. the disappearance of the blue colour, It no cuanda occura see "Secondary amines" (page 206). to boiling and add a drop or R.G. of O.S. Heat 2 o.c. of Fobling's soln, (equal volumes of No. 1 and No. 7)

cid aq. soln.) ppt. of oxalate on adding to ag. oxalic acid. The salts yield an which yields a ppt. of hydrochloride on adding to did HCL and w its bydrochloride are here considered. As usually met with pheny-the state and a yealow or brownish only liquid or crystalings mass. a hydrazine derivative (Only phenylhydrazine C.H. WH. NH. and

Bi .8.0 11 denyfhydrazone, m.p. 153"-156", as follows :--Confirm the identity of the base by preparing benzaldebyde

and shake. of O.S., heat itst to boiling, and 5 drops of bentaldebides (a) the free base, to 2 c.o. of 50% acetic acid add 5 drops

water. Heat until solution is complete, then add a equal bulk of crystallised sodium acetate and b co. (b) a hydrochloride, to 1-in, layer of O.S. in a t.t. add an eto, as indicated below. Add 5 o.c. water, shake vigorously and cool. Pilett,

er on filter paper over a small flame, and determine the mpratallise the phonylhydrazone twice from alcohol, dry a thin Filter off the solid phenylbydrazone and wash it with cold water. drops of benzaldehyde, shake vigorously and ood.

. Na No. on a soin. of O.S. consists of an insolution and and particle. Dit Jib ye motion out and be the motion of dil. IICI Secondary amines

edt bles bas feamt gaitereque lieme e otal noislume adt mo - swollol as enimasoriin a lo sonward adi mnino enima Vashooses a mori ben equivalent of 1-in, layer in a t.t. of urea (to destroy excess of HNO.).

41

Leave the funnel unstoppered, shake round and allow to stand for 5 min. Add 5 c.c. ether, shake and allow to stand until two welldefined layers are formed. Run off the lower layer, and wash the ethereal soln. with successive quantities of water (3 c.c.) until the lower layer, after running off, gives no blue colour on the addition of starch iodide soln. Separate as much water as possible from the ethereal soln. Pour 2 drops of the latter through the neck of the funnel into a dry t.t. and add 5 drops cone. H.SO., A deep blue colour indicates that O.S. is diphenylamine or one of its salts; see under "O.S. solid" (page 208). If no blue colour is obtained add R.G. of phenol and rotate the tube to mix the contents. A green or blue green colour, turned violet-red with a drop of water, and blue on adding aq. NaOH until the mixture is alkaline. indicates a nitrosamine. (Liebermann's reaction.)

Identification of O.S.

Follow the appropriate procedure, i.e. under "O.S. liquid" (below) or under "O.S. solid" (page 207).

O.S. liquid,

(Methylaniline, b.p. 192°, and ethylaniline, b p. 206°, only are here considered.)

To I c.c. of O.S. add 2 c.c. scetic anhydride and one drop of cone. H.SO, and allow to stand 5 min. Add 10 c.c. water, then add cone. NH OH until the liquid is just alkaline, cool and stir. If (a) solid separates, filter it off, dissolve it in 10 c.c. boiling

5% acetic acid: cool. Filter off the crystals, dry, and determine the m p.

M.p. 101° indicates that O.S. is methylandine. C.H. NH CH.,

(The acctyl derivative of ethylaniline melts at 54° and when prepared in the above manner usually remains as an oil.) Confirm the identity of O.S. by the preparation of the p-toluenesulphonyl derivative, m.p. 94" (see page 218).

(6) no solid separates, to I c.c. of O.S. add 10 c.c. of a saturated soln, of piene seid in benzene and stir. Filter, wash the solid picrate carefully with benzene, crystallise from aqueous alcohol, dry, and determine the m.p.

Mp. 137° indicates that O.S. is ethylamine,

#### C.H. NH-EL

O.S. solid.

If the eq. soln, of O.S. is definitely scal, proceed as under B (page 209), otherwise as under A (page 208).

Aid R.C. of O.S. to I c.c. cold cone. II. 50, and shake, Aid R.C. of C.S. dissolves 5 ir bling a deep yellow soln.

Carbazolo. (Insoluble in the HCL)

result wall be obtained.

Othersoryl derivative, m.p. 191° (see page 217).

aq. soln. strongly alkaline.)
Apply Simon's teet for a soluble secondary amine
(page 213), using 2 R.O. of O.S., when a positive

(Forms a hydrate, m p. 44°. Readily soluble in water!

104° Fiperazine, VH CH<sub>1</sub>CH

Age and deformable the Age (Allo properties). (The properties (Personal Personal Per

Allow the confents of the flack to Joyness 20 of the Mark The Real gradies and the Conference of the Mark The M

20 b. 3 (10 c. of 10 c. of 10

Derimities of diplompte.

To the soln, add one drop of 21% aq. NaXO.

Diphenylamine. (C,II,), Mil.
Add R.G. of O.S. to I c.c. cold conc. H,SO, and the

file sage ess.) \*101 .q.m. p. 107\* (see page 217).

Monacle derivative, m.p. 107\* (see page 217).

21. Denzylanijno, C.H.-NH CU. C.H., (Hydroklatik sp. M.p.

A. Aq. soin. of O.S. not acid.
Apply the test given, or prepare the derivative indicated and a state of the secondary amine whose m.p. is identical with a

A' C' II' [O' HYTOGEA]

쁘

To the soln, add one drop of 21% aq. NaNO,

—deep green colour.

Picrate (orango), m.p. 182\* (see page 53).

B. Aq. soln. of O.S. definitely acid.

A salt of an aromatic secondary amine is indicated. (Only habde salts, sulphates, and oxalates of methylanilme, ethylanilme, benzylamline, and diphenylamine are here considered.)

If no acid radical has been detected apply the following test:— Dissolve R.G. of O.S. in 2 c.c. boiling 60% acctic acid and add 2-3 drops aq. CaCl. A white ppt. indicates an exalate.

Identification of the amine.

4)

In the case of a diphenylamine salt the amine will have been already detected; diphenylamine, owing to tax weak basic character, may be blorasted by boiling the salt with water, and its identity rasy be confirmed by folliening it off, washing with water, daying, and determining the m.p. (547). In the case of a salt of any other amine, isolate the base in the manner described under "Salta of aromatic primary monamines" (page 198) and identify it in the manner indicated under "O.S. bullu" (page 207) or under "O.S. solid "(page 207).

## S KOLLOJE

DEUE, OR RED-BROWN COLOUR WITH DIL. COMPOUNDS YIELDING A GREEN, VIOLET

D- N-0X (sonozonos) Andionics Andionics solonis soloni

(a) Dissolve R.G. Of O.S. in 2 o.c. water. To the old solls had be defined the definition of the colour colour.

(b) Dissolve R.G. of O.S. in 2 o.c. water. Add the old solution old solution of the colour of the c

to 2 c.c. Br. water, to 2 c.c. water, and to 2 c.c. Br. water, to 2 c.c. Br. water, or per per colonical particles of the colonical colonical water, and the colonical colonical

Wide-talow, with evolution of gas, becoming yolow on health.

Fraction (Pyramidou)

O— V-M

O— V-M

Ph.N.— C.N.(Mo), m.p. 107°-109°.

C.N.(Mo), m.p. 107°-109°.

(a) To j.in. layer of O.S. in a c.t. add 2 c.o. dil. H.SO., warm

Pre Squitton is complete, then done to the 2 drops seq. [FeC].

TeCl. — violet-blue colour.

Tecl. Discover. To the cold solar and solar s

bad 2 o.e. aq. AgWo., does y bloke coloured are about § min.; the soluthen becomes openes owing to the formation of a grey ppl. of Ag.

Red-Brown paper or consum.

After the mixture has stood for at least 5 min said 10 e.e. sq.

N. M. II.

II. (a) the mixture is dark brown, see "m-Diamines".

(b) a vellow-green ppt, is obtained, cool, add 3 c.c. ether. oork the tube and invert two or three times.

A green ethereal layer indicates that a p-nitroso-com-

pound is present. See "Dialkylanilines" (page 211). m-Diamines, Im-Phenylenediamine; m-Tolylenediamine. Col-

ourless when pure; darken rapidly in the air.)

If the ag. soln. of O.S. is not acid, a free base is indicated; identify by m p. and by the preparation of a derivative (see list of m-diamines below).

If the aq. soln. of O.S. is definitely acid, a salt is indicated (hydrochlorides and sulphates only are considered). For identification of the base prepare a benzoyl derivative (see page 217).

NH. 63° m-Phenylenediatnine

Dibenzovl derivative, m p.

Discetyl, m p. 191°.

CH. NH. 99° m-Tolylenediamine, Dibenzovl derivative, m.p. Ňн.

224°. Discetyl, m.p. 224°.

(For preparation of derivatives, see page 217.) Dialkylanilines (Dimethylaniline, b.p. 192°, diethylapiline, b.p.

213°). O.S. liquid.

To fin. layer of powdered oxalic acid in a t.t. add 5 c.c. alcohol, heat until solution is complete, then cool. Add I c.c. of O.S. and shake.

If a white ppt is formed in a few seconds, filter, wash the solid with sloobel, dry, and determine the m.p. M.p. 139" indicates that the ppt, is the said exalate of dimethylaniline, C.H. N(CH.),

If no ppt is obtained, add I c.o. of O.S. to 5 c.c. of a saturated soln, of pierie acid in benzene, cool, and scrape the glass in contact with the liquid with a glass rod in order to assist crystallmation. Filter, carefully wash the solid with alcohol, crystallies from alcohol, dry, and determine the m.p. M.p. 142° indicates that the picrate is that of diethylaniline, C.H. N(Et).

O.S. solid, Aq. soln, definitely acid.

A salt is indicated. If no acid radical has been detected, apply the following test :- Dissolve R.G. of O.S. in 2 c.e. boiling 50%

Stoling App stuly A . + IDaD .ps sqorb C-2 lebs lats lates of erre N' C' IL [O, HALOGEN]

solution is complete, cool and your the soln, into a spanish To abrity the base, boil 2 g. of O.S. with 10 ca. dd Hillmin . SIMINAGO MA

Cool, and scrape the glass in contact with the fiquid with a feet in the flack 5 c.c. of a saturated soln, of pictic and in lemma cther (see page 21). Distil off the ether and add to the realist lumbel, Aid 10 co. of 20% ag, KOH and extract with 10 c.

tol in order to assist crystallisation. Filter, carefully was its

acid picture with alcohol, crystallise from alcohol, dry, and

.. damethylaniliae. sulfinelydists to ties a si , 8.0 tedt setseibni \*211 .q.16

.091 "

213

ij

#### SECTION 6

### ALIPHATIC AND HETEROCYCLIC AMINES

Procedure for the identification of O.S.:-

Determine the bp. and refer to the following appropriate last (its. according to whether C.S. is miscale with water, or sinks in water) of bp s of amines. If one of these bp s is identical with, or near to, that of O.S., confirm the identity of O.S. by applying to it any tests given, and by preparing and determining the mp of one or more of the derivatives there undeated. For methods of preparation of derivatives (if not given under the amino), see page 215.

O.S. miscible with water.

55° Diethylamine. (CaHa), NH. Ammoniacal, fishy odour.

Simon's test for a secondary amine.

Prepare a soln, of acetalichysic as foliows: Widone end of a piece of stoat copyer ware, about 81 an long, ant turnes round a glass rod so as to form a close spiral. Hold the straight end of the wire by means of tongs, and heat the spiral in a finem until the metal becomes constit with the black that the straight of the black of the straight of the about 10 miles of the straight of the straight of about 10 miles with 10 miles with 10 miles and 10 miles about contained in a t.t. Withdraw the spiral, and the lought, and repeat the present strate.

Add one drop of O.S. to 2 oc. of 1% aq. sodium introprusade, then add 1 oc. of the freshly prenared sola, of acetallebrile.

—deep blue colour develops in a few secs. p-Toluenosulphon)1 derivative, m.p. 60° ss-Nitro-

prodensulphonyl derivative, in p. 60° services betweenlightnyl derivative, in p. 66°. Compound with phenyl socyanate, in p. 65°.

102. Disease Cit Cit Nil Temperated miles

ant colour. (Pumes in the air, owing to the formation of carbinate.)

(page a post in the author attention and sovice

B.p. Quinoline.  $\bigoplus_{M}$  Characteristic odour.

O.S. sinks in water. Di.p. toluenesulphonyl derivative, m.p. 100°.

TITLE DO ODSOTAGO (N) sag lo notiniove na nedw (851 egaq) & tesT viqqA (Distinction from pyridine.) Gives no ppt, with aq. HgCl, or Mayor's resease.

110. Ethylenediamine, CH. NH. Ammoniacal odour.

Picrate, m.p. 164". the tube, odour very disagreeable.

h solis eds no soembnoo bus eliteib lie nwordsteed has HOM biles to D.H bbA

(urb 1113) Schoidsen enibirty to seem enillaterry a surror clear yellow liquid is obtained, which, on cooling Warm slightly-vigorous reaction occurs, and a methyl jodide,—soln, becomes yellow and cloudy

(c) Into a dry t.t. pour one drop each of 0.5, buds. bus a on mana solarages bas guitest no seviossib and at the gast out to agorb a tuode to nothibbe permanent, pale yellow ppt, is obtained after the Mayer's resgent (see page 244), doup by dop be not, EO to Gorb end one did, HCl add one drop of (6)

furning ambited ppt. of pyridine metal-4DgH .Ps a.o 2 ot .2O lo qorb a bbA (a)

(.enibradiq mon mining), and a moing at moloo suld on sevici

115° Pyridine. (Varient odour. Compound with phenyl isocranate, mp 1717.

"High other was m. p. 93" Math, mp 191 P. Toluenesulphonyl derivative, mp. 100°, Berme boniatio ed live threen salimie

Apply Sumon's test under "Diethylamine," what

- (a) To 2 c.c. dil. HCl add one drop of O.S., then add one drop of Mayer's reagent (see page 244), —immediate, pale yellow ppt., which dissolves on heating and separates again on cooling.
- (b) To 2 c.c. dil. HCl add one drop of O.S., then add one drop of a soln, of iodine in KI, —inmediate, red-brown ppt.
- (c) Preparation of quinoline tertrate, m.p. 126°.
  To 1-in. Layer of powdered tertario seid in a t.t. add 5 c. water, and shake until solution is complete. Add 1 c.c. of O.S. and shake or stir until solid separates. Filter, wash the solid with alcohol, crystallise from a small quantity of water,
- dry, and determine the m.p. (d) Preparation of quincline dichromate. (Orangeyellow.) To 1-in. layer of powdered K, Cr, O, in a t.t. add

5 c.c. dil. HCl, heat until soln. is complete, then cool.

Add 4 c.c. of O.S. and shake,

—orange ppt. (The salt may be recrystallised

from water, m.p. about 105°. Darkens near m.p.)
Picrate, m.p. 203°.

TRODS OF PREPARATION OF THE DERIVATIVES INDICATED IN THE FOREGOING LISTS OF AMINES

See sections on "Crystallisation," and "Drying of substances," ges 16-21.)

Toluenesulphonyl derivatives.

See page 218.

exenesulphonyl, or m-nitrobenzenesulphonyl derivatives. Follow the procedure given on page 218 for the preparation of columnsulphonyl derivatives, using the appropriate sulphonyl logids.

impounds with phenyl irocyanate.

To a soin, of \( \frac{1}{2} \) e.c. of plenyl isocyanate in 10 c.c. dry petroleum ser add, drop by drop, \( \frac{1}{2} \) e.c. of O.S. Filter off the white ppt, ich is formed immediately, and wash it well with petroleum ser, Dry, and determine the m.p.

crates,

To 10 oc. of a cold asturated soln, of pieric acid in benzene add ea, of O.S. and sit. Filter and wash the solid carefully with mene. Crystalline pyridine and quincline pierates from alcohol distribution pierate is repairably aduble in alcohol and also in bensel, and piperidine pierate from water.

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Saits of sliphatic and heterocyclic amines-

beneated at motor trial a vince on membraly also enimaly filliable is taken at a vince of the same of Trilow. (With ammonia to Jellow colour a produced; all discount interest in the mount of the tube paper coloured intensity wells: dimitworksored a mission of the solid in a Li. dissolid a c. a cheest 4:2 h alos silododa na dire issusisiom need and delide to isse at loger live in the live a guide politica a soll of filter page 43. at the following tree :- of in fayer of 20 in a fee all and the the tree in the fact of th ten erung a garan yd ainomme mort benteingaisile at yam disamb to sads tota commenton visually bisaddian as to similarin ones presentationinistal, libity odours, The rate ent the achiticing to any NeOH the amines are liberated, its best

Follow the procedure given for secondary amines under also see under II. Identify the amine, if possible, in the manner described made J.

If (a) a p-toluenesulphonyl derivative is obtained, P. Tolucnesulphonyl derivatives " (page 218).

\*\* of the state of the state of dichylamists of dichylamists .q.14

(cH\*) HN\*(\*HO) 79° indicates that C.S. is a salt of dimethylamint, (C'H\*)\*NH (P.D. 22.).

(5) no derivative is obtained, follow the procedure for chenc), b.p. 105°. \*001 outlinedig to state a si S.O saids establishin \*001

benietde si evitaviroù l'anonquiseneulot-q a (i) it bumery amines.

dw

smmo C'H' MH' (p b 19.1 62° indicates that O.S. is a salt of othy-

amino, CH. WH, (gas). 77° indicates that O.S. is a salt of meth!

diamine (b.p. 116"). 160" indicates that O.S. is a salt of othykine

very tisky odour; tricthy lemino is a liquid, methylamine is an inflammable gas with a oniona Vinitrot a 10 slas a si E.O. (u) no derivative is obtained, this indicates that

oly natrocamine (for detection proceed as indicated under as bloky semima vachuoses seemats, angentin to comessentello salt in dil. HCl, primary amines yield an alcohol with a rapid B. On the addition of a conc. aq. soin, of NaNO, to a soin, of the (. 69 .q.a

"Secondary amines," page 206). With tertiary amines there is no apparent reaction.

General Methods of Preparation of Derivatives of Primary

AND SECONDARY AMINES (See sections on "Crystallisation," and "Drying of substances,"

pages 16-21).

Acetyl derivatives of primary amines. (Special methods for the preparation of acetyl derivatives of secondary amines are given under the individual amine.)

To 1 c.c. or 1 g. of O.S. in a t.t. add 2 c.c. acetic anhydride; if 0.8. is solid, warm and shake until solution is complete. Cool, pour into 10 c.c. water, and stir until the oil solidifies. Filter, and wash the solid with cold water. Add the solid to 20-30 c.c. beiling water contained in a beaker and stir; if the solid does not dissolve completely gradually add glacial acetic acid with stirring until olution is complete. (Alcohol is a more suitable solvent for acetyl lerivatives of the naphthylamines.) Cool, filter off the crystals, lry, and determine the m.p.

If p.s of acetyl derivatives of solid aromatic primary monamines. (For m p.s of acetyl derivatives of liquid aromatic primary monamines see page 187.)

lalogen absent. Mp.

1279 Acetyl p-anisidine. 1320

B-naphthylamine. 148\* n-toluidine

1650 22 22

m-nitroaniline (m-Nitroacetanilide). 1590 \*\* a-naphthylamine. 219\* p-nitroaniline (p-Nitroacetanilide). ..

I present.

178° Acetyl p-chloroaniline (p-Chloroacetanilide). r present

167\* Acetyl p-bromoanume (p-Bromoacetanilide).

f.p.s of actlyl derivatives of liquid aromatic second a mines. Acetyl ethylaniline (Ethylacetanilide).

101° Acetyl methylaniline (Methylacetanilide). enzoyl derivatives.

(a) From a free amine.

In a 100-e s. conical flack dissolve I c.c. or I g. of O.S. in

5 c.c. acctone. (See note.) Add 2 c.c. benzoyl chloride, then add 50 e.c. aq. NaOlI (the first 10 c.c. or so gradually with cooling and shaking, then the remainder all at once); "he share here herewhered as and a linearities 150

The Act her removes the minimum 4:2

The Management

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210 Markeness CH. NO. Emeralment 602

Habben absent.

Tests 50 My year of the second s

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the man in the control of the second the product of Lea and asset yes it of white party to 2.0 the party and 10 and 11 and 20 the state and to con H and to and to linearly to the months added a solutions and bed standards and which is the Harm and to hingly it and additional and a said continued and the said and an anticontain and the said continued as - 2.0 b ministrate at al material

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## MUNIO, AND AND AND THE COMPONE SECTION 7

221

71 NITRO, AZO AND AZOXY COMPOUNDS

CI present.

B.p. UNO. 245° e-Nitrochlorobenzene.

See under "O.S. solid," m p. 32° (page 229).

O.S. solid.

Italogen absent. M.p.

C.H.N:O

364 Azoxybenzene.

C.H.N

Apply Test 8B (page 171) when a positive result will be obtained, owing to reduction of O.S. to antine.

OH Yellow. Tarry odour. (a) To 1-in, layer of O.S. in a t.t. add 2 c.c. aq.

NaOH and shake for 1 min .- deep red soln. (b) Reduction to o-aminophenol, identified by its benzoyl derivative, mp. 182°.

In a 100-c.c. braker place I g. of O.S., 10 c.c. aq. NaOH and 20 c.c. water. Heat to boiling, and add solid sodium hydrosulphite gradually until the soln, becomes colouries or pale yellow. Filter and rook Filter off the crystals of e-ammorphenol, wash them with cold water, and dry. Prepare the benzoyl derivative in the manner described on page 217, and determine ite mp.

(a-Nutrophenyl bensuite, mp. 55".)

CHO

64° e-Natrobenzaklakyda, NO, Pale yellow

Apply Tret (a) under "a Narephend." No change of cultur corner in the cold, but a redbrown min so chearend after buing for I must Phrey hydraume (deep red), m n. 151". Unition -- reintermer and mp 167".

ыİ

90, se Dinitrobenseno.

wiolot-blue colour, turned violot-red by said add one drop at, NaOH, (a) Dissolve R.O. of O.S. in 2 o.c. coll section and (HOAZ Padira gailed no tuoles nword lost a servic)

Dinitrotolucne " (b) (page 200), and after filter Heduca in the manner described under (b) Reduction to m-nitroantine, mp. 114". ACK!

water. Dry, and determine the m.p. End at seillelaymen bas biles edt the rathill remove more pptd, sulphur, and cool the filtrate semilor eds to a toda of awoh lice to remove potd. sulphur, add 100 cc. water, and

VII-CO-CH3

92° o-Nitroacetanilide. ( )NOs Yellow.

in To 1-in. layer of O.S. in a t.t. add 2 c.c. pq

Wilroacetanilido " (page 228). M. p. 71°, in the manner described under "P. Julinsoning bas bine outdoe of seylonbyH (d) The solution of the solution of the solutions of the solutions of the solution NaOH and heat to boiling,

neotio acid. nowover, will not interfere with the test for anilino passing over with the steam; the colour, The distillate will be yellow, owing to o-nitro

96° m-Mitrophenol. (NO, Yellow.

.nios bor-ognero-NaOII and abake, (a) To 1-in. layer of O.S. in a t.t. add 2 c.c. my.

Ponzoste, m.p. 55". Add 2 drops aq. FeCl.,-violet-rod colour. to boiling with shaking, then cool. (b) To 5 o.c. water in a 1.t. add R.G. of O.S., beat

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7}
       NITRO, AZO AND AZOXY COMPOUNDS
   М.р.
                         CHO
```

p-Nitrobenzaldehyde Very pale vellow.

(Yields a deep red-brown soln, on boiling with aq.

NaOH.) Phenylhydrazone (red), in p. 156°.

Oxidation -- p.nitrobenzoie acid, m p. 238°.

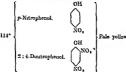
NH.

Golden vellow. 114° m-Nitroantline.

> (a) Apply the test under "o-Nitroaniline" (page 223), when a similar result will be obtained.

> (b) To 1-in. layer of OS. in a t.t. add an equal bulk of zinc dust and 5 e.c. dil. HCl Allow to stand for 5 min , shaking periodically, then filter, To the filtrate add 2 c.c. of 21% ag, NaNO. -red-brown ppt., formed from the m-phonylenediamine obtained by reduction of O.S.

Acetyl derivative, m.p. 155°. Benzoyl derivative, m.p. 155°.



(Both yield an interne yellow colour with ag. NaOH.) To 5 c.c. water in a t.t. add R.G. of O.S. heat to boiling with shaking, then cool. Add 2 drops aq. FeCl. A violet-rod colour is obtained with p-nitrophenol but not with 2:4-dinitrophenol. Acetyl derivative of p-nitrophenol, m.p. 81°.

" 2:4-dmitrophenol, m p. 72°. Benzoyl " p-nitrophenol, m.p. 142°. " 2:4-dinitrophenol, m.p. 132°.

ON 4.14 и' с' II' (0' HYTOGEN) Ш

(HO.N. ps dire gamed no suoloo aword-bot a sovid) NO. Very pale yellow. .enszendoninid-m \*00

(b) Reduction to m-nitroantline, m.p. 114°. werd. wiolet-blue colour, turned violet-red by actio Ald one drop art. NaOII, (u) Dissolve R. G. of O.S. in 2 c.c. cold acctons and

boil down to about I of the volume. Filter to to romove pptd. sulphur, add 100 c.c. water, and Dinitrotoluene " (b) (page 23), and after filtering Heduce in the manner described under "2:4

Fulter off the solid and recrystallies it from remove more potd, sulphur, and cool the filtrate.

water. Drv. and determine the m.p.

М.р.

7]

10 min. Disconnect the flask, attach it to a aloping condenser and distil until 7-8 c.c. of distillate is obtained. Test the distillate for the presence of acetic acid in the manner described on page 68, when a positive result will be obtained. To the residue in the flask add 20 c.c. aq. NaOH and shake round. Cool, filter off the nitroamine, wash it with cold water, crystallise from water, dry, and determine the m p.

COOH 238° p-Nitrobenzolo acid. Colourless.

> Apply the test under "m-Nitrobenzoic acid" (page 226) when a similar result will be obtained. Amide, m p. 201°. Methyl ester, m.p. 96°.

Ethyl ester, m.p. 57°. p-Nitrobenzyl ester, m.p. 168°. CI present.

MD.

32° o-Nitrochlorobenzene 2:4-Dinitrochlorobenzene, m.p. 50°.

44° m-Nitrochlorobenzer

Benzoyl m-chlorosniline, m p. 120°.

ca 50° 2:4-Dinitrochlorobenzer

> (Gives an orange colour on boiling with sq. NaOH.) (a) Dusoive R.G; of O.S. in 2 c c. cold acctone and add one drop aq. NaOH, -riolet-blue colour, turned riolet-red with

acetic acid. (b) Hudrolusis to 2: 4-dinitrophenol, m.p. 114".

In a 100-c.c. wide-mouthed flask place I g. of O.S., 10 c.c. ad, NaOH and 10 c.c. water. Fit

oed alob bodinom-shiw .o.o.001 a nI Hydrolysis to acclic. due to liberated p-nitroaniline.) mirw Saillod no tuoloo woller a sovid) 212° p-Nitroacetanilide. Pale yes NH-CO-CH\* ester, m.p. 93". Amide, m.p. 183°. Methyl ester, m.p. 107°. Ethi onisals and ad. MaOH, drop by drop, until the solar it and suctions bloo 2.0 2 at 2.0 to D.H evicesid award-ber a gaiblort HOaN .ps bloo ni seriossid) YON 4. 3: 5-Dinitrobenzoio acid. Pale Jellow HO00 (ce eged ess) Actions 2: 4-diminshinghlighted and sold dulute alcohol, dry, and determine the mp. receiver. Filter off the solid, organishes it from and the sold there to rather in reschedence to denote they mod if though of brouber and and hash out at harpid to samilor adt latan flask, attach it to a sloping condenser and flast the Disconnect the Disconnect the condenset, heat the contents to boing ad acme porous pot, Fit the flack with a rich Co e.c. of aq. CuSO, (Febling's No. 1 sols) and a lon-c.c. wile mouthed that place is a d05. I represent of m-destrobourse, mp 90°. 'ox decemb. 194\* 3: 4 Danteophen By Charmo N br "HN HN A' C' H' [O' HYTOCEN] 122

ш

mos bas Zailtod of stastace ad! pot. Lit the tinck with a 10 c.c. conc. HCL, 5 c.c. water

7] NITRO, AZO AND AZOXY COMPUUNDS
M.p. CH. Br

99\* p-Nitrobenzyl bromide. No.

í

p-Nitrobenzyl phenyl ether, m p. 91°.
 p-Nitrobenzyl benzoate, m.p. 89°.
 Oxidation → p-nitrobenzoic acid, m.p. 238°.

Oxidation -> p-nitrobenzoic acid, m.p. 233°.

Br

P-Nitrobromobenzene

NO<sub>4</sub>

126° p-Nitrobromobenzene

NO<sub>4</sub>

2: 1-Duutrobromobenzene, m p 72\*.

METHODS OF PREPARATION OF THE DERIVATIVES INDICATED UNDER THE NITRO COMPOUNDS IN THE FOREGOING LISTS

(See sections on "Crystallisation," and "Drying of substances," pages 16-21).

Phenylhydrazones of nitrobenzaldehydes.

Proceed as indicated under "Phenylhydrazones, O.S. solid" (145° 65).

Oxldation.

Nutrobenzaldehydes - nutrobenzose acide,

Follow the procedure under "Annakishyde," "Oxidation to anise acid "(page 61), using 1 g of O.S. Crystallise the ortho and meta compounds from water, and the para compound from alcohol.

Nutrativenes and nutrobencyl halides -- nutrobencous acula.

In a 100-ca, whe mouthed flash pixe 1 g, of O.S., 2 g, shall KHO0, 40 ca, water, 3 drups on Nolls and some process per-Dit the flash with a reduc confencer, and led the contents ratid the purple valuer of the KHnO, has designeed. Cost, three, and socilly the fitness with some IRO. Filter off the soil and wash it with ordit water. Crystillies the soil from water as the case of orthos and most composable, or from alloads in the case of the para composable; of two alloads in the case of the para composable; of two

In Desirations ( 2: Edit in Elimberta ) 2: Edition from the foreign.

Proceed as indicated under "Treparation of medicated reserve" (page 110). If O.S. is made use 1 of it.

Thenyl ether, m.p. to" (see page cyloropensono ., (e) (laca Troccol as indicated under : 2 or eistenbyll (4) -Tiolor Lor Joloir -MOAN .ps qotb ene bbs (a) Dissolve R.C. of O.S. in 2 a.c. cold Cives a yellow colour on bolling will. 72. 2:4-Dinitrobromodomsene 2: 4-m .enexnodomordominid.4: 2 41. o-Mitrobromondonino 'd'w ur present. 2: 4-Dinitrochlorobenzene, m.p. 50°. 83° p-Nitrochlorobenzene. Pale relien.

Oxidation - p-nitrobenzoie acid, m.p. 235.

p-Mitrobenzyl benzoste, m.p. 89. p-Mitrobenzyl phenyl ether, mp. 91".

71. p-Nitrobenzyl chloride.

ದಿ,ಟ್ Phenyi ether, m.p. 69" (see page 53).

the circula dil HCL dry, and determine the come, HCl and oool. Filter off the sold, oriely the section of the sould down; alm Il al galliod suminon bas gailiod of stantaco ed ted members ander a dilw stead edd N' C' H' [O' HYPOGEN]

475

CCE

Very pale yellow.

ssolve it on warming. Add a drop of phenolphthalein soln , then id 20% aq. KOH, drop by drop, until a permanent red colour is st obtained. Add just sufficient of the carboxylic acid to remove e red colour. Evaporate to dryness on a water bath, and to e residue add 1 g. p-nitrobenzyl bromide (or chloride), 21 c.c. ater and 5 c.c. alcohol. Fit the flask with a reflux condenser, at the contents to boiling, and continue boiling for 1 hr. (If imping is caused by the separation of solid, add more alcohol.) ol, and if necessary add water to precipitate the ester. Filter f the solid, wash it well with cold water, crystallise from alcohol, y, and determine the m.p.

reparation of a benzoyl derivative of a primary amine

formed by reduction of a nitro compound.

In a 100-c.c. flask dissolve (if necessary by warming) 1 c.c. or 3. of O.S. in 20 c.c. alcohol. Pour in 5 c.c. conc. HCl, then add all quantities of zinc dust, shaking round after each addition, til the liquid is practically colourless. Allow the mixture to tle, then decant off the liquid into a 250 c.c. separating funnel, ol, pour in 2 c.c. of benzoyl chloride and then add 50 c.c. aq. OH. Cool, and shake vigorously for 10 min. Extract the nzoyl derivative with about 30 cc. ether (see page 21). Wash othereal soln., first with dil. HCl, and then with water. Distil the other, crystallise the residue from alcohol, dry, and determine · ш.р.

### ADDITIONAL NITRO-COMPOUNDS

ggestions for the identification of O.S.

For the identification of alkyl esters of mtrocarboxylic acids proceed as indicated under A (page 107). The m.p. of the corresponding acid is given under the name of the ester.

To distinguish between mono-, di-, and trinitro compounds dissolve R.C. of OS. in 2 c.c. cold acctone and add one drop sq. NaOH. A blue colour indicates a dintro compound, and a red colour indicates a trinitro compound; mononitro compounds do not give a colour. (R. W. Bost and F. Nicholson, Ind. Eng. Chem. Anal., 1935, 7, 190.)

For reactions and derivatives of the compounds consult the various works of reference.

logen absent.

iquids.

B.D.

265° c-Nitroanisole, NO<sub>2</sub>-C<sub>4</sub>H<sub>4</sub> OMe. 267° c-Nitrophenetole, NO, C.H. OEt. and the second and the discount of the second and t

Proceed as indicated major "Ambies, smitten, at easter (lage 50) using, of course, sthyl should I easter. p-Nitrobenayl phenyl siber.

from dilute acetic acid. Amides and esters of nitrocarboxylic acids.

owing to its more regular descrite.
If O.S. does not dissalve resulty in the reagnit, suns usual for O.S. does not dissalve resulty in the reagnit, suns usual usual or of the control of

ing in with the state of the st

Monominities.
In a 100-or, which make place J g. of 0.3 and 10 an

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(a) of nitrophonols.

Proceed as indicated under "Acstates" (page 52).

Acetyl derlyatives.

4 al (20 egs) "seasoned" rabino bedseini se beccord kerani (O. s.Y. es seu lonedqouintal 2: 4-dinitrophened use sq. ver introphened to a solid schibides (no reliants elsecated the control Holax alica standa for the control interpretation of the control of th

is solid use I g. of it. Benzoyl derivatives of nitrophenols and nitroamines.

2: 4-Dinitrotoluene. Proceed as indicated under "Toluene" (page 116). If 02. M.p. 201\* p-Nitrobenzamide. NO.-C.H.-CONH.

(p-Nitrobenzoio acid, m.p. 238°.)

218° 3.Nitrophthalic acid. NO<sub>2</sub>C<sub>2</sub>H<sub>2</sub>(COOH), (3:1:2).
230° 5.Nitrosalicylic acid. NO<sub>2</sub>C<sub>2</sub>H<sub>2</sub>(OH)-COOH (5:2:1).
237° c.Nitrocinnamic acid. NO<sub>2</sub>C<sub>2</sub>H<sub>2</sub>CH:CH:CH-COOH.

285 P-Nitrocinnamic acid. NO<sub>8</sub>-C<sub>8</sub>H<sub>4</sub>-CH : CH-COOH.

Chlorine present.

46° o-Nitrobenzyl chloride. NO₂'C,H₂'CH₂'Cl.
Oridation→o-nitrobenzole acid, m.p. 147°.

Oxidation→o-nitrobenzoie acid, m.p. 147°.

54° Nitro-p-dichlorobenzone. NO<sub>1</sub>-C<sub>2</sub>H<sub>1</sub>-Cl<sub>2</sub> (2:1:4).

83° Picryl chloride. Yellow. (NO<sub>3</sub>), C<sub>2</sub>H<sub>2</sub>-Cl (2:4:6:1).

ernos

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.89

.02

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43.

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p-Nikobenzanilde, NO.C.H.NH.Dz. Piccamids. Yellow. (NO.), C. II, MII, (2:4:6:1). (e.Nitrobenzemide, NO.C.H.-CONH, (e.Nitrobenzemide, NO.C.H.-CONH, 2; 4.Dintrobenzelo scid, (NO.), C.H.-COOH, NO., C.H. COOH, NO.

> p.Dimitrobonzene, (NO.), C.H. (7:1:9:F)

3: 5-Dinitrosalicylio soid. (NO.),C.H. (OH) COOH.

. Picramio acid, Brownish red, (NO,),C,H,(OH)-WH. (p.Nitrocianamio acid, m.p. 255°.) . Mothyl p-mitrochmannate, NO. C.H. CH. CH. CH. COOMs on-Mittopenzanning NO.C.H. NH. Bz. p-Mikrophonylacetic soid, NO.C.H. CH. COOH.

(pt-Mitrobenzolo acid, m.p. 141°,) 2º m-Witzobenzamide, NO C.H. CONH. (P-Nitrocuntatino acid, m.p. 285".) 8. Ethyl p-nitrocinnamate. NO.C.H.CH: CH-CODEL 11. 2; 4 Dinitroacetamilde. (NO.), C.H. NHAs. 96. Methyl p-nitrobenzonte, NO.-C.H., COOMe. (p-Mitrobenzonte acid, m.p. 238°.) 94° c. Nitrobenzanilide, Yellow, NO.C.H. NHBz. (P-Nitrobenzoio soid, m.p. 238°.) 93. b-Mittopeuxyl eloohol. NO.C.H.CHOH. 88 2; + Dinitroanisole. (MO.), C.H. OMe. 86 2; + Dinitrophenetole. (NO.) C.C.H. OEE. (m-Nitrobenzoia seid, m.p. 141°.) 13. Nethyl usuitobeneses NO.C.H. COONS. (o-Mitrobenzolo scid, m.p. 147") 14. o Micropentyl sloopol. NO. C.H. CH,OR. (o-Mitrocumentic scid, m.p. 237") 72. Methyl o-nitrocannemete. NO. C.H. CH : CH-COORe. 2:4:6-Trinitroanisolo. (NO,), C.H. OMe.

p-Nitrophenetola, NO, C.H. OEL

(p-Mitrobenzolo soid, m.p. 238".) 57. Ethyl p-Nitrobenzoste, NO.C.H. COOEt. p-Nuttoenisolo, NO, C,II, OM,

(o-Nitrocunnamia acid, m.p. 237°.) Ethyl o-nitrodunamete, NO. C.H. CH : CH. CH.

(w. Mitrobenzoio scid, m.p. 141".) 27 . W. Wittohenzyl sloopol, NO. C.H. CHOOL

(1:2:9:2)

ΑŢÎ

N' C' H' 10' HYTOCKAL

#### DERIVATIVES OF ALDEHYDES AND KETONES 237

B. Derivatives of benzaldehyde.

(a) To ½ in, layer of O.S. in a t., add 2 o.c. dil. H<sub>2</sub>SO<sub>2</sub>, beat to boiling and continue boiling for 15 sec. Dulute with water to 5 o.c., cool and filter. To the filtrate add an equal volume of 2½° aq. NaNO<sub>2</sub>. Add 2 o.c. of this mixture to alkaline β-napthol (¼-n. layer of β-naphhol in a t.t. dissolved in 5 o.c. aq. NaOH). If no red ppt. is obtained apply Test (b)

An immediate, red ppt. indicates that O.S. is a benzal derivative of an aromatic primary amine (— NH, in nucleus).

e.g. benzalanilme. C.H. N : CH C.H., m.p. 54°.

To identify the amine, pour 10 c.c. dl. HCl into a 100-c.c. beaker and heat to boiling. Add 2 g. of O.S. and continue boiling with string for I min., then cool and filter. Add to the filtrate 10 c.c. of 20% aq. KOH, proceeding as described under "Salts of aromatic primary monamines" (peer 1981).

(b) To 1-in. layer of O.S. in a t.t. add 2 c.e. dil. H.50, beat to boiling and continue boiling with shaking for 1 min. Cool, dilute with water to 5 c.e. and filter Add to the filtrate aq. NaOH until it is just alkaline, heat to boiling and add 1 c. Bebling seed (come) women of No 1 and No 21.

add 1 c.c. Fehling's soln. (equal volumes of No. 1 and No. 2).

Immediate reduction (i.e., blue colour disappears and orange ppt. formed) indicates that O.S. is one of the

following:—

α Benzaldorime | C.H. CH M

α Benzaldorime | 33

Ν'ΟΗ

β-Benzaldorime | 125

HON

Bonzaldehyde semicarbazone C<sub>4</sub>H<sub>4</sub>·CH: N·NH·CO·NH<sub>4</sub>.

C. Oximes and semicarbazones.

(These compounds will have been detected by Test 8.4 (page 170).

214\*

Procedure for the identification of O.S.:-

Determine the m p. and refer to the following list of m p.s of oximes and semicarbaxones. If one of these m p.s is identical with, or near to, that of O.S., confirm the identity of O.S. by following the given instructions.

M.p. Acetoxime. (CH<sub>4</sub>),C: NOH.

CH<sub>4</sub> C: NOH.

CH<sub>4</sub> C: NOH.

CH<sub>4</sub> C: NOH.



DERIVATIVES OF ALDEHYDES AND KETONES 239

-wine red colour, turned blue by acetic acid.

 (ii) From the residue prepare a phenylhydrazone in the manner described on page 65.

(Acetophenone phenylhydrazone turns brown and shrinks at 100°, and is completely melted at 102°– 103°.)

110° Piperonaldoxime. CH O CH NOH.

Hydrolyse O.S. by the method indicated under "M.p. 59° (b)." From the residue obtained after distilling off the ether prepare a phenylhydrazone in the manner described on page 66.

(Piperonal phenylhydrazone, m.p. 100°.)

Methyl ethyl ketone semicarbazone.

CH C: N NH CO NH

Proceed as indicated under "M.p. 59" (a)" when similar results will be obtained, except that in Test (i) the wine red colour is little affected by actic acid.

(2:4-Dinitrophenylhydrazone of methyl ethyl ketone, m p. 111\*.)

141 Benzophenoxime. (C.H.), C: NOH.
Hydrolyse O.S. by the method in

Hydrolyse O.S. by the method indicated under "Mp. 55" (b)." From the residue obtained after distilling off the ether prepare a phenylhydrazone in the manner described on page 65.

(Benzophenone phenylhydrazone, m.p. 137°.)
Acctone semicarbazone. (CH<sub>4</sub>)<sub>4</sub>C: N.NH-CO-NH<sub>4</sub>.

Proceed as indicated under "M.p. 59° (a)," when similar results will be obtained.

193\* Acetophenone semicarbazone.

Proceed as indicated under "Mp. 53" (b)," when similar results will be obtained.

CH. C: NOH

(a) Dissolve R.G. of O.S. in 2 a.c. sloobel and said one



Reduction test for nitriles.

To 5 c.c. alcohol in a t.t. add one drop of O.S. and a piece of lean sodium (roughly 1 in. cube). When all the metal has disapseared add one or two drops of chloroform and heat to boiling in fume cupboard. -obnoxious carbylamine odour, indicating that primary amine has been formed. Immediately the odour is letected, cool, and add excess of cone. HCl in order to destroy

reparation of an amide from a nitrile.

Into a 100-c.c. conical flask pour 10 c.c. H<sub>1</sub>O<sub>1</sub> (15-20 volume) nd 2 c.c. aq. NaOH. Add 1 c.c. of O.S. (or 1 g. if solid) and stand he flask in water which has been heated to about 40°. (Do not ork the flask as oxygen will be evolved) Periodically remove be flask and shake vigorously until all the oil has disappeared, and aly a white solid is present. Filter off the solid, wash it with old water, crystallise from water, dry, and determine the m.p.

(Benzontrile and phenylacetonitrile are converted to the amides about 10 min.; the tolunitriles require from one to several

## ALKYL NITRITES AND NITRATES

Apply both the following tests to O.S.

(a) Into a dry t t. pour one drop of O.S. and add one drop of cone. H, SO4. The immediate evolution of red fumes indicates that O.S. is a nitrite. (The liquid also becomes red-brown in colour.) With alkyl nitrates no red fumes are evolved and the liquid remains colourless.

(b) To 5 c.c. of a cold mixture of equal volumes of water and cone, H,SO, add a trace of diphenylamine, shake for 1 min. (see note 1), then add one drop of O.S. An immediate, deep blue colour indicates that OS. is a nitrite,

Determine the bp. of O.S. and refer to the following list of b p.s of alkyl nitrates (see note 3).

If a deep blue colour is not immediately obtained, or only a pale blue colour is produced (see note 2), stand the t.t. in boiling water for 1 min., then remove and shake. A deep blue colour indicates that O.S. is a nitrate. Determine the b.p. of O.S. and refer to the following list of bps of alkyl nitrates (see note 3).

Norrs.

(1) A pale blue colour may be obtained when the diphenylamme is added to the acid, owing to the presence in the

128.	38, N F		NO CA	1 .811	oinloT-q
		shnomia.	Cell, CN	121.	oiosans@
_FIE	180.	Pites that of	Toluntinle Cy		outoT-m
.15	.01Z <sup>1</sup>	n water. Odour resembling	1112		
,17[	.102	elduloenI	CN almimioT.e	105,	oiuloT-9
			Phenylacetonitrile	40.	Phenylacetio
"Tå!	221.	more to	Propionitries Pleasan	_	Propionio 4
_	.86				oissoA
	.18	management STATE N			betsetab bis A.
p d K	(B		defermining the b.I. (ebilor tracesupile)	Vd 10 6 ATS 6	m.p. or man

se attended to bodiem tol) obinus gainbrogermon of the qua Solution the identity of S.O. by preparing and determining the general test for nitriles.

below. The reduction test (described on page 241) is a confirmatory by hydrolysis (Test 9 B, page 171), See list of carboxylio adds Districtly of O.S. is indicated by the carboxylic acid produced

### NILBITER

determine the m.p. Filter, crystallise the solid from water, dry, and complete. Cool, add 10 c.c. water and shale, sectio ambydrade. Heat until solution is just and a dry t.t. place | g. of O.S. and atty the a nl (b) Preparation of discelyl derivative, m.p. 115. -red ppt. of nickel dimethylglyoxime. (R.C. of solid dissolved in 2 c.c. water.)

ΑIJ

drop dil. NH,OH. Add to nickel sulphate soln. N' C' H' [O' HYTOGEN]

Ot Z

(2) I drop Dragendorff's (Thresh's) reagent.

-orange red ppt.

(Thresh's reagent :- Dissolve 8 g. Bi(NO.), in 20 c.c. HNO. [] pt. conc. HNO., 2 pts. water); also dissolve 27 g. KI in 40 c.c. water. Mir the solutions with constant stirring and allow to stand. Pour off from the crystals of KNO, which separate, and make up to 100 c.c. with distilled water.)

(3) I drop of a soln, of jodine in KI. -brown pot.

(4) 10 drops of naturated ac, pierie said.

--- vollow not. Nore

Salts of pyridme and of quinoline give a ppt, with Mayer's reagent If on treatment of O.S. with aq. NaOH (Test 1, page 166) only drops were obtained, isolate the base in the manner described under "Salts of aromatic primary monamines " (page 198), then proceed as indicated under "Quincline," (c) and (d) page 215. If with the aq. NaOH no oily drops were obtained, but a pungent, poculiar odour was produced, see under "Pyridine" (page 214).

SCHEME FOR THE DETECTION OF THE POLLOWING ALKALOIDS OR THEIR SALTS :- Quinine, quinidine, cinchonine, cinchonidine, morphine, codeine, anomorphine, diamorphine (heroin), strychnine, brucine, atropine, cocaine.

A. Place R.G. of O.S. in a t.t., add 2 drops dil. II SO, dilute with water to 10 cc. and look through the depth of the liquid over a dark surfam.

If a blue fluorescence is observed apply to the soin, the tests for quinino and quindine (page 246); if pegative results are obtained apply Test (c) under B for einchonidine, since this alkaloid yields a slight blue fluorescence.

If there is no blue fluorescence apply Test B.

B. Moisten a trace of O.S. with cone, HNO.

Colour produced.

-purrie red, changing to red-brown indicates that O.S. as apomorphine or one of its salts. Apply the ourfirmatory tests (page 247).

-immediate blood-red, indicates that O.S. is brucine (m.p. 1767 or one of its salts. Confirm as follows :-Evaporate the red liquid to dryness on a water bath. Add a drop of SpCl, and and star, - racket colour. (The SoCl, soln, may be prepared by discovers bin, layer of sold Soll, in a t.t. in 2 c.c. bot conc. HCl. then coving and diluters with 2 c.c. water.)

FUIOTYYTY

and siles that to chickalla renommon out to each out agreed.

n' c' n' lo' nytogeni

ome alkaloids will remmenly be encountered either as the tree (,consistential is an elicabilital substance,) amogmos sait takt homiolni need even yldeslorq lile tore

oid on the addition to their eq. soln, of eq. NaOH or eq. con eat to Aqq a bloir bas oldolos plianen era atles oil sar misser in eldulos vigariaqa vino to elduloani era abiolasia erit et or as a sait, others only as saits.

ly be met with either as the free base, or as the hydrochlorids inidine, cinchonine, cinchonidine, brucine and occaine will Iy in excess of aq. NaOH but not in excess of aq. Na,Oo. M. (Morphine, owing to its phenolic character, dissolved

present se tatrate; atropine se salicylate; and strychme eddition to the free base, bydrochloride, or sulphate, morphine chlorido. phate; diamorphine and apomorphine neually only as the

cine will usually be present as free base or phosphate; and rate, phosphate, or acetate.

to, formate, acetate, lactate, citrate, and saleylate. es hydrobromide, hydriodide, phosphate, hypophosphute, e as free base, sulphate, or hydrochlorids, and is also met

out to second edit evourer base himpel out mont it wented by ... anda to fitte the specified reagent to the depth of about moisten a substance with a reagent, dip the rounded end of only to an ordinary pin-head. larps vidguor constadue to tanome na batesibni ei cont a CONCERNING ARE LESSES.

evad lilw street evitieoq e estes est lo eno to biolaste na et. tests for alkaloids and their saits. with the moistened end of the glass rod, a neek of the bottle; then stir the substance, in a porceland ent by touching the end of the glass rod against the shade

A bas E siso Tosotions under Tests 3 and 4. bentitve result should be obtained. 2 sof Vigga enantedus labiolaula an ei BO tedt and hined in Test I (below), See note, (page 245).

diagest esting 2 o c. dil, HCl, then add the stated number of drops of biolasia son lo D.H eviosab slast gaiwolloi ed lo eno

w wieder a toolous.

ΔIJ

4

I drop Dragendorff's (Thresh's) reagent.
 —orange red ppt.

-orange red ppt.

(Thresh's reagent:-Dissolve 8 g. Bi(NO<sub>4</sub>), in 20 c.c. HNO<sub>4</sub>

(1 pt. conc. HNO<sub>2</sub>, 2 pts. water); also dissolve 27 g. KI in 40 cc. water. Mix the solutions with constant sturing and allow to stand. Pour off from the crystals of KNO<sub>2</sub> which separate, and make up to 100 cc. with distilled water.)

separate, and make up to 100 c.c. with (3) I drop of a solo, of iodine in KI.

-brown ppt.

(4) 10 drops of saturated aq. picric acid,

-yellow ppt.

Norz.

Salts of pyridine and of quinoline give a ppt. with Mayer's reagent.

If on treatment of 0.S. with aq. NaOH (Test 1, page 163) only drops
were obtained, isolate the base in the manner described under
"Salts of arounds primary monamines" (tage 168), then proceed
as indicated under "Quinolines," (c) and (d) page 215. If with the
q. NaOH no oily drops were obtained, but a pungent, pecular
q. NaOH no oily drops were obtained, but a pungent, pecular

odour was produced, see under "Pyrdine" (page 214).
Scheme for the Detection of the following Alexandre or
The Salts Salts :-Quinine, quinkline, cinchonine, cinchonidine, mor-

TRIES SALTS :—Quinine, quinkline, cinchonine, cinchonidine, morphine, codeine, apomorphine, damorphine (heroin), atrychnine, brucne, atropine, occaine.

A. Place R.G. of O.S. in a s.t., add 2 drops dd. H.SO., ddute with

vator to 10 c.c. and look through the depth of the liquid over
a dark surface.

If a blue fluorence is observed apply to the soln, the trets for so (page 248); if prigative results are

-bribe bbr (cocsine beimsuffensie). eq. ICMnO, and shake, -clear yellow soin. Add one drop cone, 11Cl, -yellow ppi (b) Dissolve It.O. of O.S. or the residue in 1 c.c. water, add 1 c.e.

(a) Dissolve R.G. of O.B. or the residue in 1 a.a. aq. Kaliba oubien odt ot (6) bna (a) atsoT Lique be .Q. with two or three drops of dil. HCl to dryness on a mater lat

11 O.S. in a sale apply Toute (a) and (b) ; if a fron alkaloid sepole ocures urb 88.

spangies indicates that O.S. is cinchonins.

Continue shaking for about I min. The separation of gold Warm slightly and shake, - ppt, dissolves.

-pale yellow flocoulone ppe. "ujo

Dissolve R.O. of O.S. in 2 drops dil. HCl, add I a.c. K. FelC. Cinchonine, m.p. 265°, and cinchonidine, m.p. 210°, changes to violet.

deep yellow colour. Add a drop of alcoholiu KOII,-col "ON"N Stir R.O. of O.S. with 2 drops cone, H.O., and a trace of Atropine, m.p. 116".

-deep violet colour changing to red. MaO, and stir,

(b) Moisten a trace of O.S. with conc. H.SO., aid a trace of change to red takes place more quickly on warmut.

-deep bine colour changing to violet, then to red. (The vanadate dissolved in 100 c,c. conc. H,50.h. (a) Moiston a trace of O.S. with Mandelin's resent a metaiold (b) Strychnine, (,enieboo to O.S. is morphine. (No colour is obtained with diamorphine that esteoibai (quorg autocadq & ot sub) motos suld qeeb A To the soin, obtained add one drop aq. Fells. water bath. Dissolve the residue in I c.c. water. or three drops dil. HCl and ersporate to depute on a

248

#### SCHEME V

Compounds containing N and S (not as sulphate)

If O.S. is a liquid see "Isothiocyanates" (page 252); if a solid follow the procedure below.

O.S. solid.

If O.S. contains

(a) a metal, proceed as indicated under "Metal present" (page 250).

(b) no metal, follow the procedure below.

No metal present.
(1) To i.in. layer of O.S. in a t.t. add 2 cc. aq. NaOH, gently

shake and hold a narrow strip of moistened red litimus paper in the mouth of the tube, taking care not to touch the glass. If the colour of the paper is (a) changed to a definite blue (inducting that O.S. is an ammonium sait), add R.G. of O.S. to 2 cc. water in a

ammonium sait), add R.G. of O.S. to 2 c.c. water in a t.t. and shake until solution is complete. Add 2 c.c. aq. FcC<sub>1</sub>, and fill up the t.t. with water. A deep red colour indicates that O.S. is a thioryanate.

If a deep red colour is not produced, proceed as industed under Test 2 in order to ascertain if O.S. is an amino-sulphonate . 'chonate; if it is neither, proceed in the colour order and in a cale of hydrocarbons

motrote of the LL

Interest to the control of the contr

(Let 1 age) (page 17.1) (All inclusions of the continuity of the c

ventrol poisson, values (e.go. 2717; general poisson variety of land variety of land variety of land to squit-stein of jill of 18 at 26 bill o

1841. "OR, H. son says, the ke all 5 though some Holes, the control of the card of the car

(page 254). (b) unaflocted, or little affected, apply Test &

If the colour of the paper is (a) rhanged to a definite blue (the colour of MH, will use ally be detected) see "Sulphonamides, and sacchain

when she no sobs possibly at all other late. I (HOAR, per all the condition of the forestable of the condition of the conditi

AMINOSULPHONIC ACIDS AND THEIR SALTS 251

(page 249) in order to ascertain if O.S. is an aminosulphonate, a mirrosulphonate, or the sodium derivative of saccharin (soluble

### AMINOSULPHONIC ACIDS AND THEIR SALTS

(Only sulphanilic, metanilic, and naphthionic acids and their alkali salts are here considered. The free acids decompose on heating.)

Procedure for the identification of O.S.:-

Apply the following distinguishing tests. Also apply any tests given under the name of the aminosulphonic acid indicated, and confirm its identity (if a free acid) by a determination of its equiv. wt. by Method 1 (page 81).

Distinguishing tests.

(1) To jim layer of O.S. in a t.t. add 2 c.c. aq. Na<sub>2</sub>CO<sub>2</sub> (or 2 c.c. water if O.S. is an alkali salt), warm until solution is complete, then fill up the t.t. with water.

If a violet fluorescence is observed see "Naphthionic acid"; if there is no fluorescence apply Test 2 in order to distinguish

between sulphanilio acid and metaniho acid.

(2) To 2 c.c. water add R.G. of O.S. and heat until solution is complete. To the hot soln add Br water until after shaking.

(a) 10 2 c.c. water add R.G. of O.S. and next until solution is complete. To the hot soin, add Br water until after shaking, the soin, is pale yellow. If a ppt, is obtained see "Sulphanite acid," or if there is no ppt. see "Metanilie acad."

Sulphambo acal Equiv. wt.

Equiv. wt. (anhydrous) 173 1.

Yields smiline on heating with solar-lime (Tret 3, page 6).
(e) To § in layer of O.S. in a b.t. a.ld 2 c.c. dichromate mixture and heat to boiling.

-pungent colour of p-benzoquinous.
(b) Preparation of 2:4:5-relevancesches, m.p. 115°.

To a solm of O.S. in hot water add strong Be solm, (10 ea. Br. 15 g. KBr. 100 cas water) until after strong the liquid is pale policy. Filter of the pps, formed, wash it was with call water, dry, and determine the m.p. who so still in some amount or more its description with the solution in the state of the state of the solution is been proportionable to the solution of the state of the solution of the

### KYLVKYAOOIIILOÑI

M p. of sulphonimide. Minosulphonin ucid indicated. 50,

". L. O periodistribit sit rot structured of the control of the rot of the ro

# NITER SHELL GAL SCIDS AND THEIR SHEIS

Gives no characteristic octour on heating with motabules in the continuation of the co

Metanilic acid. Aguiv. #£. 173-L. H. OSO. H. OSO.

poge 6). To fin. layer of 0.2, is a t.t. add 2 c.c. dichremats rate of the dilang, —pungents odour of crasphitaquinons.

Yields a nephthylamine on heating with sold and the Land

Waphthionic acid.

N' S' C' H' [0]

(6) To 5 c.c. slochol add one drop of O.S., 1 c.c. dd HCl and the equivalent of j-in, layer in a t.t. of zinc dust. Shake, more or less continuously, for 5 min., —lock-like odour, due to the formation of thioformaldehyde,

-leck-like odour, due to the formation of thioformaldehyde, CH.S.

Filter, and to the filtrate add solid NaOH until alkaline,

then add 1-2 drops of chloroform. Heat to boiling in a

fume exploard and continue boiling for 15 sec.

—obnoxious carbylamine odour, indicating that 0 S has
been reduced to a pramary amino. Immediately the odour
is detected, cool and add excess of cone. HCl in order to
destroy the isocyanide.

(c) Preparation of a thiourea.
To 1 c.c. of O.S. in a porcelain dish add 10 c.c conc. NH<sub>4</sub>OH and evaporate on a water bath. Crystallise the solid from

water, dry, and determine the m.p.

M p. of theores. Theorets indicated.
74° Allylthoures Allyl isothicoyanate, b.p. 150°.
CH.; CH-CH.; N : CS.
Phenylthioures Phenyl isothicoyanate, b.p. 221°.

#### THIOUREAS

C.H. N . CS.

Indicated by a brown or black colour obtained after boiling with aq. NaOH and adding aq. lead acetate (Test 1, page 249).

Procedure for the identification of O.S.:-

Determine the m.p. and refer to the following list of m p.s of thoursas. If one of these m p.s is identical with, or near to, that of O.S., confirm the identity of O.S. by applying the given tests.

M.p. 74° Allylthiourea (Thiosinamine). CH<sub>3</sub>: CH-CH<sub>2</sub>:NH-CS-NH<sub>3</sub>. Preparation of acetyl derivative, m.p. 98°-100°.

To him layer of O.S. in a t.t. add 5 c.e. acetone and shake until solution is complete. To the solution add gradually 2 c.e. acetyl chioride and shake. Filter, wash the solid with acetone, dry, and determine the mp.

151° Diphenylthiourea (Thiocarbanilide).

C.H. NH-CS-NH-C.H.
Practically insoluble in hot water.

154° Phonylthiourea. C.H. NH-CS-NH., Soluble in hot water.
(a) To R.G. of O.S. in a dry t.t. add 5 drops of conc.

4HN

Hquiv. wt. 173-L.

Domino at lonningomina-su Given me characteristic odour on health with sales and H\*OS

ELLER RIBHT GNA EGION. SUNDHALUEGATIN Yields no quinone on heating with thichromate mixture.

d sad no balanch 1 same of all obimenbilging a craquel Procedure for the identification of 0.8.

and the of sulphonumble Nitronalphana be in lateral ON sobimanodqlus to a q.m to tsil gulwottol the state and , q m at orientation has (longola mort osillateyts)

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p-Toluenesulphonamide.  $\bigcirc$  SO  $_{s}$ NH,

M.p. 137°

m-Xylene-4-sulphonamide.

SO,NH,
Proceed as indicated under "Preparation of dimethyl
derivatives " (page 257). The formation of a solid
derivative, m p 80°, indicates that O S. is p-toluenesulphonamide.

With m-xylene-4-sulphonamide the oil obtained does not solidify.

SO,NH,

150° Naphthalene-1-sulphonamide.

In a 100-a.c. wide-mouthed fissis place the equivalent of jin, layer in a t. of 0.5 and add 5 c. acceled of water and cone. If SO<sub>s</sub>. Fit the fissk with a reflux condenser, beat the contents to boiling and continue boiling for 5 min. A deposit of naphthalene (characteristic dout) will be present thalene (characteristic dout) will be present water, filter it off, dry, and determine the mp. (M n. of naphhalance, 50°).

153° Benzenesulphonamide. C.H. SO NH.

154° o-Toluenesulphonamide. SO,NH,

Proceed as indicated under "Preparation of dimathly derivatives" (page 207). The dimethyl derivatives will be obtained as an oil which will not solidify on cooling and shaking. Extract the oil from the mixture with 10 co. ether (see page 21), and wash the othered soln, three times with about 3 co. water. Separate as much water as possible, distill off the other, cool the residual oil and stir

(W. epo) & set T as BE.

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or beam to this hand, and maintained
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## KINTHOOFF ORT EXCITETEOUSTOF

aux boss until schulchn is complete. To solm schi I n.s. sep. K. k'el'e(L'V)... ---green ochous, changing to blise.

-doop red soin.

[4] To 2 a.c. of this section is the lift of C. of C. and book until solution is complete. To

whill them a rayo i. 3 vin a in 20 Do. Doll sool! if a both the southern bars ables of the bill be and the southern being a single of the south in the south in the south in the south in a south in the south south south south in the south i

172. Thioures. Will, CS-Nils.

172. Thioures. Will, CS-Nils.

-180. (a) Heat R.G. of O.S. in a dry f.f. over a small thams

while all the state of some one builds all of the parts of the the state of the the parts of the the parts of 
"HOAN ps. no. 0 an heriosath us as he melamund edd os sub) melatios bor A to stadque edt tedt spiniotioni bas banoqueo uit 14 bennin sood sas entam eitsamy reamny beinded et (102 il. 2000 tijn 20 to tanantenn (4) story lydy. A stamoqueo died dire word is der lydy. A stamoqueo died dire word is der lydy. A stamoqueo died dire word is der lydy. A stamoqueo died dire

M.5.O., West for 5 sec over a small stans (100; H. o. bigh), cool, and carefully dishe with very to 5 oc. Again cool, and said 5 ce. of 25% of NaVO., Add 2 ce. of this mixture to stake for the philotophic of the mixture of stanships of the forest of graphic of the dissolved in 5 ce. aq. NaOH).

Preparation of dimethal derivatives of sulphonamides.

In a Wann, wide monthed, took place 1 2, of QS, and add 10 c.c. sq. NaOH. Heat until soln, is complete, then add to the warm soln, 2 c.o. of dimethylsulphate (see caution, page 161). Cark the flask and shake more or less continuously, for 5 min. Add another 10 c.c. of aq. NaOH, and boil under a reflux condenser for 5 min., in order to destroy the excess of dimethylsulphate. (The dimethyl derivative will be present as an oil.) Cool and shake well. If the oil solidifies, filter, wash the solid with cold water, crystallise from aqueous alcohol, dry, and determine the m.p.

### SULPHONYL DERIVATIVES OF PRIMARY AND SECONDARY AMINES

These compounds will have been indicated by Test 4 (page 250). (Only m.p.s of p-toluenesulphonyl derivatives of the commoner aromatic primary and secondary amines are here given )

The identity of OS. may be confirmed by preparing from the amine the sulphonyl derivative suspected (see page 218) and carrying out a mixed m.p. determination (see page 14).

p-Toluenesulphonul derivatives of aromatic primary ammes.

103° v-Toluenesulphonanilide

108\* p-Toluenesulphon-o-toluidide

p-Toluenesulphon-m-toluidide Soluble in aq. NaOH,

p-Toluenesulphon-p-toludide

p-Tolsenesulphonyl derivatives of aromatic secondary arrines,

M.D. 875

87° Ethyl p-toluenesulphonanilide | Insoluble in aq. NaOH.
 94° Methyl p-toluenesulphonanilide | Insoluble in aq. NaOH.

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of NCTs.

- India to the red white indicatus the present of a factorial and a

-deep red return thereby (thus to the formation of a solids as composited).

(a) the section of the first of the section and the section of t

As, N. S. Na piesent.
Sulpharenobenzene. Daolan 3:3
Stufphurephenumine. (Sulpharenobenzene. Daolan 3:3

ED THOOPY

म्पर्य के प्रस्तित के किया है किया विकास किया किया किया किया किया किया है कि किया किया किया किया किया किया कि

As, M, and Ma grename.

#### MIXTURES OF ORGANIC COMPOUNDS

Treatment of mixtures of two or more of the following:—a nstrogenous base, an acidio substance, neutral compounds.

Each constituent must be isolated and purified, and then identified in the manner described for single substances on page 3.

A few of the simpler methods of separation are described below, it being assumed that not more than one substance until be removed from the mixture by any me of the particular treatments, and that any final residue after the treatments indicated until consist of only one substance.

For details concerning extraction with ether, washing of an ethereal soln, and removal of the ether by distillation, see page 21. Also see notes (page 263).

### Procedure:-

Apply to the mixture the alkelizing test for N if this element is about no less can be pressin, and therefore the treatment of the etherat sole. With 6d MC (describer for page 259) should be omitted. If the mixture it a liquid follow the precedure given below; if a solid, proceed as indicated under "Mixture solal" (page 259). The procedure should be suitably modified if it is known that the mixture craticism only two constituents.

Misture liquid.

Four 10 &c. of the mixture into a 100 c. o, wide-mouthed flask, connect the latter to a water condenser and use all  $\gamma$ t. t. as the receiver. Heat the contents of the flask by means of a boding water bath. He nose constituent of the mixture obtails over, proceed to identify it. If the contents of the thak after cooling are (a) said, proceed as indicated under "Mixture sold!" (lage 202), and (b) the distribution of the third of the cooling are (a) of the cooling of th

as indicated under "Mixture solid" (b).
(u) a lower layer of liquid is present, run it off into a small

separating funnel, add 10 c.c. dry ether (in order to remove any of the other constituents of the mixture), shake and

seared and design than the Est with the litt and and, and the the

5 G the mount attential notes, talted nature seaton such this Intel of the other met abself the restinal beat.

william of the time, only and the handles and me Welet, office friends of, gives into mare then a slight butbally ween all libre beine ledlebult filler ribe forteite auf fine W Shaka, seperate and report the truck tayor.

mejaratung feininal anul acht 10-13 e a. uther. (a) an est or emulating, pour the contents of the leader line a

n Kimmer

has ofth filter is off, wash is well with only water, dry, and he soletaque sond e 11 THE PROPERTY OF THE PARTY OF

thin sociate sits of a major and a major thing the sits in the last after the form the sits in the sit soldaffe al onless and them until the MON .ps 202 LbA - (Derablemon ered ere sotew ni eldulos

Usultaje to sidulosni ena drida swad olizinora vino) mad a lo Treatment of the HCl extract, which may contain the hydrochbraie layer (HCl extract) into a beater. way and the mist chemical era started bendab-tlew own

insert the atopper, shake well, and allow to stand until Jill ith as off this found edt ni mice isordie edt o't -: SMOTIOI

(5) the solid has dissolved completely in the other, proceed as with a mineral soid, a sulphonic soid, an amide hydrate, a polyhydrio alcohol, a salt of an organic base

odies a the olitation a metallic sait, a carbo indicated under (b). Identify the solid substance, which the filtrate into the separating funnel, and proceed a allow the washings to fall into the main filtrate. Four order to remove the other constituents of the mixium, to Finally wash the solid in the filter with dry ether, if by washing it out with small quantities of dry ether.

Transfer any solid remaining in the flund to the Elle contents of the funnel through the neck into a filter. II (a) some solid remains undissolved, shake round and pour th

separating funnel, add 30 cc. dry ether and shake well. Introduce about 5 g. of the finely powdered solid into a 100-ci Mixiure solid.

indicated under "Mixture solid" (b). from the small funnel into the larger one, and proceed mentify it (probably a polyhydrio alcohol). Pour the ch allow to settle. Run off the lower layer into a dry tt.

layer. (This is to remove any remaining basic constituent from the ethereal soln., and, of course, will not be necessary if no base has been isolated.)

Add to the ethereal soln. 30 c.c. of aq. NaOH, shake well, and

run off the lower layer (NaOH extract) into a beaker.

Treatment of the NaOH extract, which may contain the sodium derivative of one of the following: a carboxylio acid, a phenolic compound, a \$\tilde{\ell}\$ testonic ester, a sulphonamide, ascelaring, a sulphonyl derivative of a primary amine, a member of the purine group:

Just scidify with cone. HCl, cool and stir; if no solid separates scrape the glass in contact with the liquid with a glass rod.

If (a) a solid separates, filter it off, wash it well with cold water, dry, and identify it.

(§) no solid separates, pour the contents of the beater into a separating finnel and extract with 10-15 c.c. ether, (If the acidification produced an oil or emulsion one extraction will be sufficient; if, however, no old or emulsion was obtained at least three extractions should be carried out and the several etheract extracts added together. This is necessary since a water-soluble compound may have been theracted by the HGJ.

Wash the othereal extract several times with 3 c.e. distilled water until the wash-water, after separation, gives not more than a slight turbidity on the addition of dil HNO<sub>2</sub> and aq. AgNO<sub>2</sub>. Distil off the other and identify any residue.

Trainent of the main thereof sole. (after extraction with eq. NaOH) which may contain one of the following neutral compounds:

a hydrocarbon, an alcohol, an ester, an ether, an aldebyde, a lettone.

a nitrile, etc. (substituents such as NO, or halogen may be present in these compounds):—

Weah the othereal soin, with distilled water until the weah-water is no longer alkaline. Distil off the other and identify any residue NOTES.

(1) Owing to the fact that ether is somewhat soluble in water, it may be necessary to add more of the former from time to time. Indications of the necessity for more ether are

(a) the ethereal soln, becoming cloudy.

(i) an imperfect separation of the othered and aqueous layers. (ii) M is said in produced on the adultion to the externed size, of did. HCl, or of sq. NaOH, filter by suction the whole contents of the separating funcel. What het saids will wish other, allowing the watarings to fall into the first filtrate. Wish not the separating funced with water, poor the filtrate into it and run of the lower layer. Threat the adult or the saids and the be isolated by altration or by attraction with other filtres the commercial product may be unsufficiency. The baselights only a form of the product of the commercial first a mittern of 200; Koliffed, and 430 see safety poses 50, until all the solid line indexed and the solution is just set in the methyl compare.

the aldehyde or ketone would be regenerated, and could On heating the solid or lower layer with HCl or Ma, (U, see note 2), or in solution in the lower layer. subscorp tol) bilos a sa rohite ansoring guisd rottel edt would thus be converted into its bisulphite compound. An aldehyde, and in some cases a ketone, if present shaken with a saturated soin, of sodium bisulphite. (c) The othereal soin, after the various treatments might be (101 egaq) " hotosteb mosd sad lodcola ol." robus leadrosob sads of ralimis at emboome edT could be isolated by filtration or by extraction with groups), if present as alkali salt, would be blersted and ON on this brunoquoo ollonold a 3.6) constadus buthe Uderw a madw aOO thir botsmass od tdgim (a) The MaOH extract, before scidifying with cone. Hel. (5) The scheme of separation may be extended as follows: chloride, such as that of benzylaniline.)

MIXTURES OF ORGANIC COMPOUNDATION of the maker and the maker "Treatment" Treatment of the Holl struct," or under "Treatment of the Most actinet," or under "Treatment of the Most of the Holl struct," or under "Treatment of the Most of the Holl struct," or under "Treatment of the Most of the Mos

tormortion of a sparingly soluble sodium salt, such as solum simmamate, or to the formation of a sparingly soluble hydro-

#### SPECIAL REAGENTS

Alkali-sugar and alkali-zinc mixtures.

Alkali sugar mixture.

An intimate mixture of pure anhydrous sodium carbonate and one-tenth of its weight of pure sucrose.

When organic compounds are heated with this mixture, halogens are converted into sodium halides, sulphur into sodium sulphude, and nitrogen (in a limited number of cases) into sodium cyanide.

The proportion of sugar employed does not cause excessive times when the mixture is heated, yes provides sufficient earhon for the reduction of sulphate, etc., to sulphide. The final heating to redness is essential, in order to cause any such reduction, to decompose the sugar throughly, and to remove any deposit from the mouth of the tube, so that a perfectly colouriess filtrate is obtained.

### Alkali-zine mixture.

An intimate mixture of zine dust and half its weight of anhydrous sodium carbonate (ordinary commercial products).

The gradual heating of the organic substance is necessary, as otherwise negative results in the test for nitrogen may be obtained with aromatic bases and their salts and derivatives.

The alkah-zine mixture, with constituents of A.R. quality, may also be used to test for sulphur and halogens, provided that the tests for chlorine and sulphur are made in comparison with a

blank test. (Middleton.)

The test for sulphur is carried out as follows:—After the hot tube has been plunged into water, the mixture is heated to boiling and allowed to settle, and the liquid is then decanted through a filter. To the residue is the dash (containing insoluble sine sulphids if sulphur is present in the organic compound) are added about 10 c. a. of dulte hydrechlors acid, and a filter paper (upon the centre of which a drop of sodium plambite solution has been poured) is immediately placed over the dish. If sulphur is present in the organic compound, a dark brown stain, vashlo on the upper surface of the paper, will be formed.

Blank tests usually give a slight brown stain visible only on the under surface of the paper. pea chorine into cold weter until se asturated solution is qualified to the light of the cold and the cold into th

to obtained to secretary the street of bromine will discolar the solution of the street of the secretary and the secretary and s

Bromline water. Shake 5 e.e. bromine with 100 e.e. water until no mo.

Bromine in carbon terrachloride. Dissolve 4 c.c. bromine in 100 c.c. carbon tetrachlori

Dissolve 13 g. of crystallised neutral coppor acetals is of cold I.% acetic acid soln.

Brombe in carbon tetrachloride.

Soo page 214. Barloed's reagent. Dissolve 13 g. of cretallised neutral copper acetate

Alkaloidal reagents. See page 214.

the organic and series of the cyanida to change present in the cyanida as subject of throeparate bonds present in the cyanida and the cyanida

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abounded.

With picto send polynitro compounds, the pictor and polynitro and send polynitro send and polynitro specification of the class of compound the following an indication of this class of compounds.

(U) The tests are applicable without modificatin n to very concliding an experience, e.g. ethyl bronde for the passes of prints and narrated of organs (iii) Except in the case of prints and paracted of organs compounds the reactions are perfectly quiet, even with

cutting particular base requestional to to apparentable and the recent of the control of the con

## Dichromate mixture.

Dissolve 100 g. of sodium dichromate in a mixture of 250 c.c. conc. H<sub>2</sub>SO<sub>4</sub> and 750 c.c. water.

# 3:5-Dinitrobenzoyl chloride.

This reagent is employed for the conversion of sloohols and phenols into solid esters. Since the reagent decomposes on keeping under ordinary conditions, it is advisable to prepare it as required by the action of PCl<sub>2</sub> on 3:5-dinitrobenzoic acid.

# Preparation of 3:5-dinitrobenzoates of alcohols.

In a porcelain dish place 0.3 g. of 3:5-dimitrobenzois acid and add i.in. layer in a t. of powdrant PCi<sub>1</sub>. Grind the substances together with a pastic until the mixture becomes semi-solid. Add 1.e. of the abcolod and stir like mixture becomes semi-solid. Add 1.e. of the abcolod and stirring, for 5 min. if the abcolod is primary or for 30 min. if the abcolod is prompt of the 30 min. if the abcolod is prompt of the 30 min. if the abcolod is prompt of 50 min. if the 50 mi

### NOTE.

In some cases it may be necessary to add more alcohol in order to effect solution of the derivative. Derawtives of low mp. such as n-anyl 3:5-dinirobemosts, mp. 46°, should be dissolved in petroleum ether air blown, by means of bellows, over the surface of the liquid until the derivative crystallases out.

Methyl		and det	IVALIA	crys	COLLI	508 QU	t,
Ethyl	3:5-dinit	robenzo	ate.			m.p.	107°
n-Propel	**	**	•			**	93°
iso Propyl	**	**				••	73*
n-Butyl	**	••				**	122°
iso Butvi	••	**				••	64*
ecc.Butyl	**	**		•	•	**	87°
*-Amyl		,,	•	٠			76°
Cirlo Haran	. "	**					46*

## Fehling's solution.

- Dissolve 69 3 g. of pure crystallised copper sulphate in water containing a few drops of dil. H<sub>2</sub>CO<sub>2</sub> and make up to 1 litre.
   Dissolve 346 g. of Rochelle salt /sodium potassium tarteste/and
  - make up to 1 litre.
    mately equal volumes immedi-

893

Schiff's reagent.

Dilute 10 c.c. of the stock solution with water to 1 litre, add as a stock solution. SO, until no further change occurs. Filter, and keep the filtrate Dissolve I g. of a resentitine salt in 100 c.c. warm water. Pass

Sodium nitrite solution (21%). overnight. 50 c.c. of a saturated aqueous soln, of SO, and allow to stand

Sodium nitroprusside solution (1%). Dissolve 12-5 g, of MaNO, in 500 a.c. water,

kocping, becoming green. Dissolve 0-1 g. in 20 o.c. water. The solution deteriorates on

Approx. 2N. Acids (dilute) and alkalies (aqueous). Common reagents.

Lerrie chloride (freshly prepared). Industrial spirit unless absolute alcohol is specified. JodoolA.

5 g. of hydrated salt dissolved in water and made up to

Other reagents as used for Inorgania Analysis. 100 c.c.

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WITH SUGGESTED EXERCISES FOR STUDENTS WHO DO NOT EXQUIRE IDENTIFICATION WORK

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- 4. Clear and systematic tables have been drawn to take various observations.
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